

WORLD INTELLECTUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:		(11) International Publication Number: WO 99/32514
C07K 14/00	A2	(43) International Publication Date: 1 July 1999 (01.07.99)
(21) International Application Number: PCT/US (22) International Filing Date: 15 December 1998 (30) Priority Data: 60/068,179 19 December 1997 (19.12. 60/099,840 11 September 1998 (11.09) (71) Applicant (for all designated States exc. WARNER-LAMBERT COMPANY (US/U Tabor Road, Morris Plains, NJ 07950 (US). (72) Inventor; and (75) Inventor/Applicant (for US only): SUN, Yi [US/Hillway Court, Ann Arbor, MI 48105 (US). (74) Agents: RYAN, M., Andrea; Warner-Lambert Cor Tabor Road, Morris Plains, NJ 07950 (US) et al.	97) 98) ept U SS]; 4	CZ, EE, GE, HR, HU, ID, IL, IS, JP, KP, KR, LC, LK, LR, LT, LV, MG, MK, MN, MX, NO, NZ, PL, RO, SG, SI, SK, SL, TR, TT, UA, US, UZ, VN, YU, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TI, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BI, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG). Published Without international search report and to be republished upon receipt of that report.

(54) Title: SAG: SENSITIVE TO APOPTOSIS GENE

(57) Abstract

The present invention provides novel genes and polypeptides derived therefrom encoding a redox-sensitive protein that promotes cell growth, protects cells from apoptosis, scavenges oxygen radicals and can be used for the reversion of a tumor phenotype. In an attempt to identify gene(s) responsible for 1,10-phenanthroline (OP)-induced apoptosis in tumor cells we have used the differential display technique and cloned an OP-inducible gene, SAG (Sensitive to Apoptosis Gene). SAG encodes a novel, redox-sensitive, heme-binding protein with a zinc RING finger domain. The SAG protein consists of 113 amino acids with a calculated molecular weight of 12.7 kDa. Sequence homology searches reveal that SAG is highly conserved among species, suggesting its functional importance. This suggestion is demonstrated by the finding that SAG disruption in yeast is lethal. Two SAG deletion mutants have been detected in human cancer cell lines originating from colon and testis, suggesting its possible role in human carcinogenesis. Overexpression of SAG protein in a human colon carcinoma line, DLD1, and a human neuroblastoma line, SY5Y, protects cells from apoptosis induced by OP, zinc and copper ions. Furthermore, antisense SAG transfection inhibits certain tumor cell phenotypes in DLD1 human cell line and microinjection of SAG RNA stimulates cell growth. We propose that SAG protein is a cellular protective molecule functioning as a redox sensor to buffer oxidative-stress induced damage as well as a growth factor to stimulate cell growth. SAG protein will be an ideal molecular target in the development of drugs against neurodegenerative disorders, cancers, muscle dystrophy, and promoting wound healing.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AL	Albania	RS	Spain	LS	Lesotho	SI	Slovenia
AM	Armenia	FI	Finland	LT	Lithuania	SK	Slovakia
AT	Austria	FR	France	LU	Luxembourg	SN	Senegal
AU	Australia	GA	Gabon	LV	Latvia	SZ	Swaziland
AZ	Azerbaijan	GB	United Kingdom	MC	Monaco	TD	Chad
BA	Bosnia and Herzegovina	GE	Georgia	MD	Republic of Moldova	TG	Togo
BB	Barbados	GH	Ghana	MG	Madagascar	TJ	Tajikistan
BE	Belgium	GN	Guinea	MK	The former Yugoslav	TM	Turkmenistan
BF	Burkina Faso	GR	Greece		Republic of Macedonia	TR	Turkey
BG	Bulgaria	HU	Hungary	ML	Mali	TT	Trinidad and Tobago
BJ	Benin	IE	Ireland	MN	Mongolia	UA	Ukraine
BR	Brazil	IL	Israel	MR	Mauritania	UG	Uganda
BY	Belarus	IS	Iceland	MW	Malawi	US	United States of America
CA	Canada	IT	Italy	MX	Mexico	UZ	Uzbekistan
CF	Central African Republic	JP	Japan	NE	Niger	VN	Viet Nam
CG	Congo	KE	Кепуа	NL	Netherlands	YU	Yugoslavia
CH	Switzerland	KG	Kyrgyzstan	NO	Norway	ZW	Zimbabwe
CI	Côte d'Ivoire	KP	Democratic People's	NZ	New Zealand		
CM	Cameroon		Republic of Korea	PL	Poland		
CN	China	KR	Republic of Korea	PT	Portugal		
CU	Cuba	KZ	Kazakstan	RO	Romania		
cz	Czech Republic	LC	Saint Lucia	RU	Russian Federation		
DE	Germany	LI	Liechtenstein	SD	Sudan		
DK	Denmark	LK	Sri Lanka	SE	Sweden		
EE	Estonia	LR	Liberia	SG	Singapore		
1							
1							

DNC ---

SAG: SENSITIVE TO APOPTOSIS GENE

Background f the Inventi n

5

10

15

20

25

30

The present invention relates to a novel gene and polypeptides derived therefrom encoding a redox-sensitive protein that protects cells from apoptosis and promotes cell growth, as well as antibodies directed against the polypeptide. The invention also describes methods for using the novel gene, polypeptides, and antibodies in the detection of genetic deletions of the gene, subcellular localization of the polypeptide, isolation of discrete classes of RNA, inhibition of apoptosis, scavenging of oxygen radicals, reversion of tumor phenotype, and therapeutic applications by gene therapy.

Summary of the Related Art

Apoptosis, also referred to as programmed cell death, is a genetically programmed process for maintaining homeostasis under physiological conditions and for responding to various stimuli (Thompson (1995) Science 267, 1456-1462). This form of cell death is characterized by cell membrane blebbing, cytoplasmic shrinkage, nuclear chromatin condensation, and DNA fragmentation (Wyllie (1980) Int. Rev. Cytol. 68, 251-306). The process of apoptosis can be divided into three distinct phases: initiation, effector molecule stimulation and DNA degradation (Kroemer et al. (1995) FASEB J. 9, 1277-1287; Vaux and Strasser (1996) Proc. Natl. Acad. Sci. USA 93, 2239-2244). Apoptosis can be initiated in various cell types by a wide variety of physical, chemical, and biological stimuli (both internal and external), including diverse cancer therapeutic drugs, oxidative DNA damage reagents, and cytokines (Kroemer (1997) Nature Med. 3, 614-620, White (1996) Genes Dev. 10, 1-15; Sen and D'Incalci (1992) FEBS Lett. 307, 122-127; Dive and Hickman (1991) Br. J. Cancer 64, 192-196; Yuan et al. (1993) Cell 75, 641-652). These initiators trigger the effector molecules in cells leading to apoptotic signal transduction and amplification, which ultimately results in irreversible DNA degradation and cell death.

Many genes are involved in the apoptotic process. In general, the products of these genes are classified as either inducers or inhibitors of apoptosis. The balance between the activities of apoptosis inducers and inhibitors in a given cell determines whether that cell undergoes apoptosis. Among the growing list of apoptotic regulatory genes, the most well characterized are the p53 tumor suppressor gene, the Bcl-2 gene family (consisting of both inducers and inhibitors of apoptosis), the interleukin 1β converting enzyme (ICE) gene family, and FAS/Fas ligand (Kroemer (1997), White (1996); Yuan et al. (1993); Nagata and

Golstein (1995) Science 267, 1449-1456). During apoptosis, there are substantial interactions involving the products of apoptotic regulatory genes, including heterodimer formation among the gene products of the Bcl-2 gene family, and p53 activation of Bax expression (Oltvai et al. (1993) Cell 74, 609-619; Miyashita and Reed (1995) Cell 80, 293-299).

The inventor has recently found that 1.10 phenanthroline ("OP"), a metal chelating agent, can activate p53 activity and induce apoptosis in two murine tumor cell lines that harbor endogenous wild-type p53 (Sun et al. (1997) Oncogene 14, 385-393). OP is a typical metal chelating reagent in that it chelates Fe(II) and prevents Fe(II)-mediated hydroxyl radical formation through the Fenton reaction (Halliwell et al. (1989) in: Free Radicals in Biology and Medicine, 2nd ed., Clarendon Press, Oxford; Auld (1988) in Methods in Enzymology, Vol. 158 (J. F. Riordan and B. L. Valle, Eds.) PP. 110-114, Academic Press, New York). OP has been shown to prevent hydroxyl radical-induced DNA damage in a number of cellular systems (Sun. Y. Free Radic, Biol. Med. 8:583-599 (1990); Martins and Meneghini, Biochem J. 299:137-140 (1994); Morgan et al., Biochem, Pharmacol, 44:215-221 (1992)). Activation of p53 by OP was found to significantly contribute to, but was not required for subsequent apoptotic cell death (Sun et al., (1997) Oncogene 14: 385-393; Sun (1997) FEBS Lett. 408, 16-20). Thus, the critical genes and gene products responsible for OP-induced apoptosis remain to be characterized. A better understanding of the molecular mechanisms of apoptotic induction will allow improved design of therapeutic drugs that either induce (anti-cancer) or inhibit (anti-aging) apoptosis.

Summary of the Invention

5

10

15

20

25

30

The present invention provides novel genes and polypeptides derived therefrom encoding a redox-sensitive protein that protects cells from apoptosis, scavenges oxygen radicals, and can be used for the reversion of a tumor phenotype.

In one aspect, the present invention provides novel isolated and purified DNA sequences (referred to herein as "mouse SAG" and "human SAG") as shown in SEQ ID 1 and SEQ ID 3, and their gene products (referred to herein as "mouse SAG protein" and "human SAG protein") as shown in SEQ ID 2 and SEQ ID 4, that are induced during 1,10-phenanthroline ("OP")-induced apoptosis. In another embodiment, the present invention comprises a nucleotide sequence that hybridizes to the nucleotide sequence shown in SEQ ID 1 and SEQ ID 3 under high stringency hybridization conditions. In a preferred embodiment, the isolated and purified DNA sequence consists essentially of the DNA sequence of SEQ ID 1 or SEQ ID 3.



In another aspect, the invention provides novel recombinant DNA molecules, comprising SAG subcloned into an extra-chromosomal vector. In a further aspect, the present invention provides recombinant host cells that are stably transfected with a recombinant DNA molecule comprising SAG subcloned into an extra-chromosomal vector.

In a different aspect, the present invention provides a substantially purified recombinant protein comprising a polypeptide substantially similar to the SAG protein shown in SEQ ID 2 and SEQ ID 4. In a further aspect, the present invention provides a polyclonal antibody that selectively binds to proteins with an amino acid sequence substantially similar to the amino acid sequence shown in SEQ ID 2 and SEQ ID 4.

5

10

15

20

25

30

Additional aspects of the present invention provide a method of detecting the SAG protein in cells, comprising contacting cells with a polyclonal antibody that recognizes the SAG protein; a method of detecting cells containing SAG deletions, comprising isolating total genomic DNA from the cell and subjecting the genomic DNA to PCR amplification using primers derived from the DNA sequence of SEQ ID 1 and SEQ ID 3; and a method of detecting cells containing SAG deletions, comprising isolating total cell RNA and subjecting the RNA to reverse transcription-PCR amplification using primers derived from the DNA sequence of SEQ ID 1 and SEQ ID 3.

In another aspect, the present invention further provides methods of isolating RNA containing stretches of polyA, polyC, or polyU residues from cells, contacting the total cell RNA with the SAG protein, and incubating the RNA-SAG protein mixture with an antibody that recognizes the SAG protein.

In another aspect of the present invention, a method for isolating genes induced during cell apoptosis is provided, comprising treating cells with OP, subjecting the OP-induced RNA to the differential display procedure, and cloning the OP-induced genes.

A further aspect of the invention provides a method for protecting mammalian and/or non-mammalian cells from apoptosis induced by redox reagents, comprising introducing into mammalian and/or non-mammalian cells an expression vector comprising a DNA sequence substantially similar to the DNA sequence shown in SEQ ID 1 and SEQ ID 3, which is operatively linked to a DNA sequence that promotes the expression of the DNA sequence, wherein the isolated and purified DNA sequence of SEQ ID 1 and SEQ ID 3 will be expressed at high levels in the mammalian and/or non-mammalian cells.

An additional aspect of the present invention provides a method for treatment of mammalian and/or non-mammalian tumor cells, comprising introducing into mammalian and/or non-mammalian tumor cells an expression vector comprising a DNA sequence

substantially similar to the DNA sequence shown in SEQ ID 1 and SEQ ID 3, which is operatively linked to a DNA sequence that promotes the expression of the antisense strand of the DNA sequence, wherein the antisense strand of the DNA sequence of SEQ ID 1 and SEQ ID 3 will be expressed at high levels in the mammalian and/or non-mammalian cells.

Another aspect of the present invention provides a method for oxygen radical scavenging in an organism, comprising administering an oxygen radical-reducing amount of a pharmaceutical composition comprising SAG protein and a pharmaceutically acceptable carrier.

A further aspect of the present invention provides for gene therapy applications of SAG, including but not limited to methods of promoting the closure (i.e., healing) of a wound in a patient.

The foregoing is not intended and should not be construed as limiting the invention in any way. All patents and publications cited herein are incorporated by reference in their entirety.

15 Brief Description of the Drawings

5

20

25

30

Figure 1A. Predicted structural features of the deduced protein sequence of the mouse and human SAG cDNA.

Figure 1B. Description of human SAG protein mutants.

Figure 2. Bar graph depiction of soft agar colony growth of various SAG-transfected stable cell lines.

Figure 3. Graphical representation of tumor mass in SCID mice per days post implant with SAG transfectants.

Detailed Description of the Invention

The present invention provides novel genes and polypeptides derived therefrom encoding a redox-sensitive protein that protects cells from apoptosis, scavenges oxygen radicals, and can be used for the reversion of a tumor phenotype. The present invention also comprises genes and their gene products involved in OP-induced apoptosis. The isolation of such genes and their gene products permits a detailed analysis of the OP-induced apoptotic pathway, thus providing laboratory tools useful to identify the mechanisms of OP-induced apoptosis and enabling improved design of therapeutic drugs to regulate apoptosis.

Within this application, unless otherwise stated, the techniques utilized may be found in any of several well-known references such as: Molecular Cloning: A Laboratory Manual

(Sambrook, et al., 1989, Cold Spring Harbor Laboratory Press), Gene Expression Technology (Methods in Enzymology, Vol. 185, edited by D. Goeddel, 1991. Academic Press, San Diego, CA), "Guide to Protein Purification" in Methods in Enzymology (M.P. Deutshcer, ed., (1990) Academic Press, Inc.); PCR Protocols: A Guide to Methods and Applications (Innis, et al. 1990. Academic Press, San Diego, CA), Culture of Animal Cells: A Manual of Basic Technique, 2nd Ed (R.I. Freshney. 1987. Liss, Inc. New York, NY), and Gene Transfer and Expression Protocols, pp. 109-128, ed. E.J. Murray, The Humana Press Inc., Clifton, N.J.).

5

10

15

20

25

30

In one aspect, the present invention provides novel isolated and purified DNA sequences, hereinafter referred to as Sensitive to Apoptosis Genes ("SAG"), encoding SAG proteins. In one embodiment, the invention comprises DNA sequences substantially similar to those shown in SEQ ID 1 (mouse SAG) or SEQ ID 2 (human SAG), respectively. As defined herein, "substantially similar" includes identical sequences, as well as deletions, substitutions or additions to a DNA, RNA or protein sequence that maintain the function of the protein product and possess similar zinc-binding motifs. Preferably, the DNA sequences according to the invention consist essentially of the DNA sequence of SEQ ID 1 or SEQ ID 3, or are selected from the group consisting of SEQ ID 11, SEQ ID 13, SEQ ID 21, SEQ ID 23, SEQ ID 25, SEQ ID 27, SEQ ID 29, SEQ ID 31, SEQ ID 33, SEQ ID 35, SEQ ID 37, SEQ ID 39, SEQ ID 41, SEQ ID 43, SEQ ID 45, SEQ ID 47 and SEQ ID 49. These novel purified and isolated DNA sequences can be used to direct expression of the SAG protein and for mutational analysis of SAG protein function.

Mutated sequences according to the invention can be identified in a routine manner by those skilled in the art using the teachings provided herein, as described in Example 8, *infra*, and techniques well known in the art.

In another embodiment, the invention comprises a nucleotide sequence that hybridizes to SEQ ID 1 and/or SEQ ID 3 under high stringency hybridization conditions. As used herein, the term "high stringency hybridization conditions" refers to hybridization at 65°C in a low salt hybridization buffer to the probe of interest at 2 x 108 cpm/µg for between about 8 hours to 24 hours, followed by washing in 1% SDS, 20 mM phosphate buffer and 1 mM EDTA at 65°C, for between about 30 minutes to 4 hours. In a preferred embodiment, the low salt hybridization buffer comprises between, 0.5-10% SDS, and 0.05M and 0.5 M sodium phosphate. In a most preferred embodiment, the low salt hybridization buffer comprises, 7% SDS, and 0.125M sodium phosphate. These DNA sequences can be used to direct expression

of the SAG protein and for mutational analysis of SAG protein function, and are isolated via hybridization as described.

5

10

15

20

25

30

In another aspect, the invention provides novel recombinant DNA molecules, comprising SAG or a sequence substantially similar to it subcloned into an extrachromosomal vector. This aspect of the invention allows for *in vitro* expression of the SAG gene, thus permitting an analysis of SAG gene regulation and SAG protein structure and function. As used herein, the term "extra-chromosomal vector" includes, but is not limited to, plasmids, bacteriophages, cosmids, retroviruses and artificial chromosomes. In a preferred embodiment, the extra-chromosomal vector comprises an expression vector that allows for SAG protein production when the recombinant DNA molecule is inserted into a host cell. Such vectors are well known in the art and include, but are not limited to, those with the T3 or T7 polymerase promoters, the SV40 promoter, the CMV promoter, or any promoter that either can direct gene expression, or that one wishes to test for the ability to direct gene expression. These recombinant vectors are produced via standard recombinant DNA protocols as described in the references cited above. This aspect of the invention allows for high level expression of the SAG protein.

In a further aspect, the present invention provides recombinant host cells that are stably transfected with a recombinant DNA molecule comprising SAG subcloned into an extra-chromosomal vector. The host cells of the present invention may be of any type, including, but not limited to, non-eukaryotic (e.g., bacterial), and eukaryotic such as fungal (e.g., yeast), plant, non-human animal, non-human mammalian (e.g., rabbit, porcine, mouse, horse) and human cells. Transfection of host cells with recombinant DNA molecules is well known in the art (Sambrook et al., Molecular Cloning, A Laboratory Manual, 2nd ed., Cold Spring Harbor Press, 1989) and, as used herein, includes, but is not limited to calcium phosphate transfection, dextran sulfate transfection, electroporation, lipofection and viral infection. This aspect of the invention allows for *in vitro* and *in vivo* expression of SAG and its gene product, thus enabling high-level expression of SAG protein, as described in Example 6, *infra*.

In another aspect, the present invention provides a substantially purified recombinant protein comprising a polypeptide substantially similar to the SAG polypeptides shown in SEQ ID 2 and SEQ ID 4. Furthermore, this aspect of the invention enables the use of SAG protein in several *in vitro* assays described below. As used herein, the term "substantially similar" includes deletions, substitutions and additions to the sequences of SEQ IDs 1-4 (as



appropriate) introduced by any *in vitro* means. As used herein, the term "substantially purified" means that the protein should be free from detectable contaminating protein, but the SAG protein may be co-purified with an interacting protein, or as an oligomer. Preferably, the protein sequences according to the invention comprise an amino acid sequence selected from the group consisting of SEQ ID 2, SEQ ID 4, SEQ ID 12, SEQ ID 14, SEQ ID 22, SEQ ID 24, SEQ ID 26, SEQ ID 28, SEQ ID 30, SEQ ID 32, SEQ ID 34, SEQ ID 36, SEQ ID 38, SEQ ID 40, SEQ ID 42, SEQ ID 44, SEQ ID 46, SEQ ID 48, and SEQ ID 50. In a most preferred embodiment, the protein sequences according to the invention comprise an amino acid sequence selected from the group consisting of SEQ ID 2 and SEQ ID 4. Mutated sequences according to the invention can be identified in a routine manner by those skilled in the art using the teachings provided herein and techniques well known in the art. This aspect of the invention provides a novel purified protein that can be used for *in vitro* assays, as described in Examples 12, *infra*, and as a component of a pharmaceutical composition for oxygen radical scavenging, described *infra*.

In a further aspect, the present invention provides antibodies and methods for detecting antibodies that selectively bind polypeptides with an amino acid sequence substantially similar to the amino acid sequence of SEQ ID 2 and SEQ ID 4. The antibody of the present invention can be a polyclonal or a monoclonal antibody, prepared by using all or part of the sequence of SEQ ID 2 or SEQ ID 4, or modified portions thereof, to elicit an immune response in a host animal according to standard techniques (Harlow and Lane (1988), eds. Antibody: A Laboratory Manual, Cold Spring Harbor Press). In a preferred embodiment, the entire polypeptide sequence of SEQ ID 2 or SEQ ID 4 is used to elicit the production of polyclonal antibodies in a host animal.

The method of detecting SAG antibodies comprises contacting cells with an antibody that recognizes SAG protein and incubating the cells in a manner that allows for detection of the SAG protein-antibody complex. Standard conditions for antibody detection of antigen can be used to accomplish this aspect of the invention (Harlow and Lane, 1988). This aspect of the invention permits the detection of SAG protein both *in vitro* and *in vivo*, as described in Examples 12 and 14, *infra*.

In a further aspect, the present invention provides a diagnostic assay for detecting cells containing SAG deletions, comprising isolating total genomic DNA from the cell and subjecting the genomic DNA to PCR amplification using primers derived from the DNA sequence of SEQ ID 1 SEQ ID 3, SEQ ID 11, SEQ ID 13, SEQ ID 21, SEQ ID 23, SEQ

ID 25, SEQ ID 27, SEQ ID 29, SEQ ID 31, SEQ ID 33, SEQ ID 35, SEQ ID 37, SEQ ID 39, SEQ ID 41, SEQ ID 43, SEQ ID 45, SEQ ID 47 and SEO ID 49.

This aspect of the invention enables the detection of SAG deletions in any type of cell, and can be used in genetic testing or as a laboratory tool. The PCR primers can be chosen in any manner that allows the amplification of a SAG gene fragment large enough to be detected by gel electrophoresis. Detection can be by any method, including, but not limited to ethidium bromide staining of agarose or polyacrylamide gels, autoradiographic detection of radio-labeled SAG gene fragments, Southern blot hybridization, and DNA sequence analysis. In a preferred embodiment, detection is accomplished by polyacrylamide gel electrophoresis, followed by DNA sequence analysis to verify the identity of the deletions. PCR conditions are routinely determined based on the length and base-content of the primers selected according to techniques well known in the art (Sambrook et al., 1989).

5

10

15

20

25

30

An additional aspect of the present invention provides a diagnostic assay for detecting cells containing SAG deletions, comprising isolating total cell RNA and subjecting the RNA to reverse transcription-PCR amplification using primers derived from the DNA sequence of SEQ ID 1 SEQ ID 3, SEQ ID 11, SEQ ID 13, SEQ ID 21, SEQ ID 23, SEQ ID 25, SEQ ID 27, SEQ ID 29, SEQ ID 31, SEQ ID 33, SEQ ID 35, SEQ ID 37, SEQ ID 39, SEQ ID 41, SEQ ID 43, SEQ ID 45, SEQ ID 47 and SEQ ID 49. This aspect of the invention enables the detection of SAG deletions in any type of cell, and can be used in genetic testing or as a laboratory tool.

Reverse transcription is routinely accomplished via standards techniques (Ausubel et al., in Current Protocols in Molecular Biology, ed. John Wiley and Sons, Inc., 1994) and PCR is accomplished as described above.

In another aspect, the present invention provides methods of isolating RNA containing stretches of polyA (adenine), polyC (cytosine) or polyU (uridine) residues, comprising contacting an RNA sample with SAG protein, incubating the RNA-SAG protein mixture with an antibody that recognizes the SAG polypeptide, isolating the antibody-SAG protein-RNA complexes, and purifying the RNA away from the antibody-SAG protein complex. This aspect of the invention provides a novel *in vitro* method for isolating a discrete class of RNA. In a preferred embodiment, the RNA sample is contacted with SAG protein in the presence (for preferential isolation of polyA and polyC-containing RNAs), or absence (for preferential isolation of polyU-containing RNAs), of a reducing agent. Preferred reducing agents for use in this aspect of the invention include, but are not limited to DTT and

β-mercaptoethanol. The reducing agents are preferably used at a concentration of between about 50 mM and 1 M. Isolation of antibody-SAG protein-RNA complexes can be accomplished via standard techniques in the art, including, but not limited to the use of Protein-A conjugated to agarose or cellulose beads.

5

10

15

20

25

30

In a further aspect of the present invention, a method for isolating genes induced during cell apoptosis is provided, comprising treating one set of cells with OP and not treating a control set of cells, isolating RNA from each set of cells, subjecting the RNA from each set of cells to reverse transcription and PCR ("differential display"), identifying cDNAs that are expressed in the OP-treated set of cells and not in the control set of cells, and cloning the OP-induced cDNAs. This aspect of the invention provides a tool for isolating other genes that control the OP-induced apoptotic pathway and is useful both as a way to enable the design of therapeutic drugs that regulate apoptosis and as a laboratory tool to identify the mechanisms of OP-induced apoptosis. Details of the differential display technique, including selection of primers, are well known in the art (Liang and Pardee, Science 257:967-971, 1992). Reverse transcription and PCR conditions are routinely determined based on the length and base-content of the primers selected according to techniques well known in the art (Sambrook et al., 1989). In a preferred embodiment, OP is used at a concentration of between 50 μM and 300 μM. In a most preferred embodiment, OP is used at a concentration of between 100 μM and 150 μM.

A further aspect of the invention provides a method for protecting mammalian and/or non-mammalian cells from apoptosis induced by redox reagents, comprising introducing into mammalian and/or non-mammalian cells an expression vector comprising a DNA sequence substantially similar to the DNA sequence shown in SEQ ID 1 or SEQ ID 3, that is operatively linked to a DNA sequence that promotes the expression of the DNA sequence and incubating the cells under conditions wherein the DNA sequence of SEQ ID 1 or SEQ ID 3 will be expressed at high levels in the mammalian and/or non-mammalian cells. In a preferred embodiment, the DNA sequence consist essentially of SEQ ID 1 or SEQ ID 3. Suitable expression vectors are as described above. In a preferred embodiment, the coding region of the human SAG gene is subcloned into an expression vector under the transcriptional control of the cytomegalovirus (CMV) promoter to allow for constitutive SAG gene expression.

An additional aspect of the present invention provides a method for inhibiting the growth of mammalian and/or non-mammalian tumor cells, comprising introducing into

mammalian and/or non-mammalian tumor cells an expression vector comprising a DNA that is antisense to a sequence substantially similar to the DNA sequence shown in SEQ ID 1 or SEQ ID 3 that is operatively linked to a DNA sequence that promotes the expression of the antisense DNA sequence. The cells are then grown under conditions wherein the antisense DNA sequence of SEQ ID 1 or SEQ ID 3 will be expressed at high levels in the mammalian and/or non-mammalian cells. In a preferred embodiment, the DNA sequence consists essentially of SEQ ID 1, SEQ ID 3, SEQ ID 11, SEQ ID 13, SEQ ID 21, SEQ ID 23, SEQ ID 25, SEQ ID 27, SEQ ID 29, SEQ ID 31, SEQ ID 33, SEQ ID 35, SEQ ID 37, SEQ ID 39, SEQ ID 41, SEQ ID 43, SEQ ID 45, SEQ ID 47 and SEQ ID 49.

5

10

15

20

25

30

In a most preferred embodiment, the DNA sequence consists essentially of SEQ ID 1 or SEQ ID 3. In a further preferred embodiment, the expression vector comprises an adenoviral vector wherein SAG cDNA is operatively linked in an antisense orientation to a cytomegalovirus (CMV) promoter to allow for constitutive expression of the SAG antisense cDNA in a host cell. In a preferred embodiment, the SAG adenoviral expression vector is introduced into mammalian tumor cells by injection into a mammalian tumor cell mass.

An additional aspect of the present invention provides a method for oxygen radical scavenging in an organism, comprising introducing into mammalian and/or non-mammalian cells an expression vector comprising a DNA sequence substantially similar to the DNA sequence shown in SEQ ID 1 or SEQ ID 3 which is operatively linked to a DNA sequence that promotes the expression of the DNA sequence, and the cells are grown under conditions wherein the DNA sequence of SEQ ID 1 or SEQ ID 3 will be expressed at high levels in the mammalian and/or non-mammalian cells. In a preferred embodiment, the DNA sequence consists essentially of SEQ ID 1 or SEQ ID 3. In a preferred embodiment, the SAG cDNA is operatively linked to a cytomegalovirus (CMV) promoter, to allow for constitutive expression of the SAG cDNA in a host cell.

Another aspect of the present invention provides pharmaceutical compositions and methods for oxygen radical scavenging in an organism, comprising administering an oxygen-reducing amount of a pharmaceutical composition comprising the SAG protein of SEQ ID 2 or SEQ ID 4 and a pharmaceutically acceptable carrier.

Chimeric gene constructs of the present invention (e.g., expression vectors) containing SAG polynucleotide sequences may be used in gene therapy applications to achieve expression of SAG or anti-sense SAG polynucleotide sequences in selected target cells, including non-eukaryotic cells (i.e., plant) and eukaryotic cells. Gene therapy applications typically involve identifying target host cells or tissues in need of the therapy,

designing vector constructs capable of expressing a desired gene product in the identified cells, and delivering the constructs to the cells in a manner that results in efficient transduction of the target cells.

The cells or tissues targeted by gene therapy are typically those that are affected by the disease that the vector construct is designed to treat. For example, in the case of cancer, the targeted tissues are malignant tumors.

5

10

15

20

25

30

In one embodiment, the present invention provides a method of promoting the closure (i.e., healing) of a wound in a patient. This method involves transferring exogenous SAG to the region of the wound whereby a product of SAG is produced in the region of the wound to promote the closure (i.e., healing) of the wound.

The present inventive method promotes closure (i.e., healing) of both external (e.g., surface) and internal wounds. Wounds to which the present inventive method is useful in promoting closure (e.g., healing) include, but are not limited to, abrasions, avulsions, blowing wounds, burn wounds, contusions, gunshot wounds, incised wounds, open wounds, penetrating wounds, perforating wounds, puncture wounds, seton wounds, stab wounds, surgical wounds, subcutaneous wounds, tangential wounds, or traumatopneic wounds. Preferably, the present inventive methods are employed to close chronic open wounds, such as non-healing external ulcers and the like.

Exogenous SAG can be introduced into the region of the wound by any appropriate means, such as, for example, those means described herein. For example, where the wound is a surface wound, SAG can be supplied exogenously by topical administration of SAG protein to the region of the wound.

Preferably, exogenous SAG is provided to the wound by transferring a vector comprising an SAG expression cassette to cells associated with the wound. Upon expression of SAG within the cells in the region of the wound, a product of SAG is produced to promote wound closure (i.e., healing). Transferring a vector comprising an SAG expression cassette to cells associated with the wound is preferred as such procedure is minimally invasive, supplies SAG products locally within the region of the wound, and requires no reapplication of salves, solutions, or other extrinsic media. Furthermore, SAG activity remains expressed during wound closure and will inactivate following healing.

The vector comprising the SAG expression cassette can be transferred to the cells associated with the wound in any manner appropriate to transfer the specific vector type to the cells, such as those methods discussed herein.

As discussed above, the cells associated with the wound to which the vector is transferred are any cells sufficiently connected with the wound such that expression of SAG within those cells promotes wound closure (i.e., healing), such as cells within the wound or cells from other sources. In one embodiment, the cells are cells of the wound, and the present inventive method comprises transfer of the vector to the cells in situ.

5

10

15

20

25

30

In other embodiments, the cells are not the cells of the wound, but can be cells in an exogenous tissue, such as a graft, or can be cells *in vitro*. For example, to promote the healing of certain types of wounds, the cells associated with the wound can be cells within a graft, such as a skin graft. Transfer of the vector to the cells associated with the wound, thus involves transferring the vector to the cells within the graft *ex vivo*. For other wounds, the cells associated with the wound are cells *in vitro*, and the cells are transferred to the region of the wound following transfer to them of a vector containing the SAG expression cassette.

The present inventive method applies to any patient having a wound. For example, the patient can be any animal, such as a mammal. Preferably, the patient is human.

In another embodiment, the present invention provides a method of inhibiting or promoting plant cell growth. The method involves the use of chimeric gene constructs to achieve expression of SAG, in the case of promoting growth of plants, or anti-sense SAG, in the case of inhibiting plants (i.e., weeds), polynucleotide sequences in selected target plant cells.

The dosage regimen for *in vivo* oxygen radical scavenging by the administration of SAG protein is based on a variety of factors, including the type of injury, the age, weight, sex, medical condition of the individual, the severity of the condition, the route of administration, and the particular compound employed. Thus, the dosage regimen may vary widely, but can be determined routinely using standard methods. In a preferred embodiment, the pharmaceutical composition comprises between 0.1 and 100 mg of SAG protein. In a most preferred embodiment, the pharmaceutical composition comprises between 1 and 10 mg of SAG protein.

The SAG protein may be made up in a solid form (including granules, powders or suppositories) or in a liquid form (e.g., solutions, suspensions, or emulsions). The SAG protein may be subjected to conventional pharmaceutical operations such as sterilization and/or may contain conventional adjuvants, such as preservatives, stabilizers, wetting agents, emulsifiers, buffers etc.

While the SAG protein can be administered as the sole active pharmaceutical agent, they can also be used in combination with one or more other agents. When administered as a

combination, the therapeutic agents can be formulated as separate compositions that are given at the same time or different times, or the therapeutic agents can be given as a single composition.

For administration, the SAG protein is ordinarily combined with one or more adjuvants appropriate for the indicated route of administration. The SAG protein may be admixed with lactose, sucrose, starch powder, cellulose esters of alkanoic acids, stearic acid, talc, magnesium stearate, magnesium oxide, sodium and calcium salts of phosphoric and sulphuric acids, acacia, gelatin, sodium alginate, polyvinylpyrrolidine, and/or polyvinyl alcohol, and tableted or encapsulated for conventional administration. Alternatively, the SAG protein may be dissolved in saline, water, polyethylene glycol, propylene glycol, carboxymethyl cellulose colloidal solutions, ethanol, corn oil, peanut oil, cottonseed oil, sesame oil, tragacanth gum, and/or various buffers. Other adjuvants and modes of administration are well known in the pharmaceutical art. The carrier or diluent may include time delay material, such as glyceryl monostearate or glyceryl distearate alone or with a wax, or other materials well known in the art.

In a preferred embodiment of the present invention, the SAG protein pharmaceutical composition is administered intramuscularly (IM) or intravenously (IV). A suitable IM or IV dose of active ingredient of SAG protein is 5 mg/mL administered daily. For IM or IV administration, the active ingredient may comprise from 0.001% to 10% w/w, e.g., from 1% to 2% by weight of the formulation, although it may comprise as much as 10% w/w, but preferably not more than 5% w/w, and more preferably from 0.1% to 1% of the formulation.

The present invention may be better understood with reference to the accompanying examples that are intended for purposes of illustration only and should not be construed to limit the scope of the invention, as defined by the claims appended hereto.

25 Examples

5

10

15

20

30

Example 1. Identification of an OP-inducible gene

The differential display (DD) technique was employed to isolate genes responsible for or associated with OP-induced apoptosis in two murine tumor lines. Since OP induced-apoptosis can be visually detected at 12 hours post exposure (Sun, (1997) FEBS Lett. 408:16-20), it was reasoned that gene(s) responsible for apoptosis induction should be up- or down-regulated prior to the appearance of apoptosis. Six hours of OP treatment was conducted, therefore, in one of these tumor lines followed by the DD analysis.

Mouse JB6 tumor line L-RT101 (an epidermal originated tumor cell line) was cultured in Minimal Essential Medium with Earle's salts (BRL) containing 5% fetal calf serum (Sigma). H-Tx cells, a spontaneously transformed mouse liver line, were cultured in Dulbecco's Modified Eagle Medium containing 10% fetal calf serum and 1 mM sodium pyruvate. Human colon carcinoma line DLD-1 was grown in 10% DMEM.

L-RT101 cells were treated with 150 uM OP for 6 hours and subjected to differential display analysis using DMSO-treated cells as a control. Briefly, total RNA was isolated from both OP-treated and control cells using RNAzol solution (Tel-Test) according to the manufacturer's instructions, and subjected to reverse transcription (RT), performed as previously described (Sun et al. (1993) Mol. Carcinogenesis 8, 49-57), followed by the polymerase chain reaction (PCR). The primer used for reverse transcription (P1) consisted of the sequence 5 AAGCTTTTTTTTTTTTTTT (SEQ ID 5), wherein R consists of either adenine, guanine or cytosine. P1 was used as the downstream primer in the subsequent PCR while the upstream primer consisted of the sequence AAGCTTNNNNNNN (SEQ ID 6), wherein N consists of adenine, cytosine, guanine, or thymine.

Primers P1 and P2 reproducibly detected differential expression between the control and OP-treated cells. The fragments reproducibly showing differential expression were PCR amplified using the same primers and used as probes for Northern analysis (Sun et al. (1992) Cancer Res. 52:1907-1915) of both L-RT101 and H-Tx cells treated with OP (Sun (1997) FEBS Letters 408:16-20). Those fragments that were induced by OP (as determined by Northern analysis) were then subcloned into TA cloning vectors (In Vitrogen) according to the manufacturer's instructions, and sequenced by DNA Sequenase Version 2.0, according to the manufacturer's instructions (Amersham). The resulting clones comprise OP-inducible cDNA fragments.

25 Example 2. cDNA library screening and 5'RACE

5

10

15

20

30

One of the OP-inducible clones was used as a probe to screen a mouse lung cDNA library to clone the full length mouse SAG cDNA. Briefly, 1 x 10^6 recombinant plaques were plated onto 1% NZY in 150 mm plates (a total of 20). The recombinant phage DNA was transferred to nitrocellulose membrane and hybridized with mouse SAG probe (2X10⁸ cpm/ μ g) in a hybridization solution containing 5X SSC, 5X Denhardt solution, 50 mM sodium phosphate, and 100 μ g/mL denatured DNA at 60°C for 16-18 hours. The filter was

then washed once for 5 min in a solution of 2XSSC/0.1% SDS, once for 5 min in 0.5XSSC/0.1% SDS, and twice 0.1XSSC/0.1% SDS for 15 min.

5

10

15

20

25

30

The longest clone isolated was a 1.0 kilobase ("kb") fragment consisting of a partial open reading frame and the entire 3'-end untranslated region. A mouse brain Marathon-Ready cDNA (ClonTech) was screened via PCR amplification using a primer derived from the 1 kb fragment and another primer derived from the vector sequence, according to the protocol supplied with the cDNA library. This yielded a further 100 bp fragment consisting of 5'-end untranslated sequence and some of the coding sequence. The derived cDNA clone consists of 1140 base pairs ("bp") (SEQ ID 1) that encode a novel deduced protein of 113 amino acids, containing 12 cysteine residues (SEQ ID 2). The open reading frame was preceded by 17 bp upstream sequence. The start codon was located in a context that conformed 100% to the Kozak consensus sequence (Kozak,M. (1991) J. Biol. Chem. 266, 19867-19870). An in-frame stop codon was identified 72 bp upstream of the start codon in the 5' untranslated region in one genomic clone (not shown). The 3'-end untranslated region consists of 792 bp sequence with two polyadenylation signals (AATAAA). These data indicate that a near full length cDNA was isolated.

The mouse cDNA was used as a probe to screen a human HeLa cell cDNA library (Strategene) as described above. One positive clone was isolated and purified through two more cycle of screening. In this manner, a 754 bp clone containing a polyadenylation signal at the 3' end was isolated (SEQ ID 3). The human cDNA also contains an open reading frame encoding a novel predicted 113 amino acid polypeptide containing 12 cysteine residues (SEQ ID 4). The sequence identity between the isolated mouse and human cDNAs is 82% in overall sequence and 94% in the coding region. At the protein level, they shared 96.5% identity, with all 12 cysteine residues being conserved. Computer analysis of protein databases using the GCG program (Genetics Computing Group, Madison, WI) revealed that the encoded proteins share 70% identity with hypothetical proteins from yeast (accession #Z74876) and C-elegans (accession #80449).

Motif searching of the deduced protein sequences using the GCG program did not reveal any known functional domains. However, they each contain two imperfect heme binding sites (CXXCH, at codons 47-51 and 50-54) (Matthews, Prog. Biophys. Mol. Biol. 45:1-56, 1985) and one imperfect C₃HC₄ zinc ring finger domain (Freemont et al., Cell 64:483-484, 1991) at the C-terminal of the molecule (Fig. 1A) among other consensus motifs. The second potential heme binding domain (Fig. 1A) contains a substitution of arginine to

histidine (amino acid 54). Since these two amino acids are structurally similar, this may constitute an authentic heme binding site. The zinc ring finger domain mismatch involves substitution of cysteine by histidine at amino acid 85. The ring finger domain in this protein is a C₃H₂C₃ structure, rather than the consensus C₃HC₄ structure. Since cysteine and histidine residues are interchangeable in zinc binding (Berg and Shi, Science 271:1081, 1996; Inouye et al., Science 278:103-106, 1997), the C₃H₂C₃ domain in these proteins may comprise authentic zinc-binding sites. Significantly, these heme and zinc ring finger domains are 100% conserved among *C. elegans*, mouse and human. In yeast, only the last cysteine residue in C₃H₂C₃ motif was not conserved. This evolutionary conservation of the heme and zinc-binding domains suggest their functional importance.

5

10

15

20

25

30

Other motifs identified in the deduced sequence of the SAG protein, when allowing for a single mismatch, include an aminoacyl-transfer RNA synthetase class II motif (codons 54-63), a Kazal serine protease inhibitor family motif (codons 85-107), a Ly-6/U-par domain (codons 65-107), a prokaryotic membrane lipoprotein lipid attachment site (codons 16-27), and somatotropin, prolactin and related hormone motifs (codons 49-66).

These experiments thus resulted in the cloning of novel mouse and human genes that encode nearly identical, evolutionarily conserved protein that contain distinct heme and zinc binding motifs.

Example 3. SAG is inducible by OP in both mouse and human tumor cells

To confirm that the cloned cDNAs are subject to OP induction, a Northern analysis was performed with RNAs isolated from mouse tumor lines L-RT101 and H-Tx, and human colon carcinoma line DLD-1. Subconfluent cells were treated with 150 μ M OP for various times up to 24 hours and subjected to total RNA isolation. Fifteen μ g of total RNA was subjected to Northern analysis using mouse SAG or human SAG cDNA as probes.

Both cloned mouse and human cDNAs detected an OP inducible transcript with a size of 1.2 kb and 0.9 kb, respectively. Since these genes were induced in the OP-induced apoptosis pathway, the genes were named Sensitive to Apoptosis Genes (hereinafter referred to as "SAG"), which encode SAG proteins.

Example 4. Tissue distribution and embryonic expression of SAG

SAG expression was next examined in multiple human tissues. The assays were performed as detailed previously (Sun et al. (1993) Mol. Carcinogenesis 8, 49-57; Sun et al., Proc. Natl. Acad. Sci. USA 90:2827-2831, 1993). Briefly, total RNA was isolated from

multiple human tissues (ClonTech) and then subjected to Northern blot analysis using the mouse or human SAG cDNA as probes. SAG RNA was detected in all tissue examined including heart, brain, pancreas, lung, liver, skeletal muscle, kidney, pancreas, spleen, thymus, prostate, testis, ovary, small intestine, colon, and peripheral blood leukocytes. A very high expression level was detected in heart, skeleton muscle and testis, which consume high levels of oxygen. Its tissue distribution and high level expression in oxygen-consuming tissues, and its induction by a redox sensitive compound (OP), implies that SAG encodes a redox sensitive protein.

5

10

15

20

25

30

Since SAG protein is evolutionarily conserved, the possible developmental role of SAG was tested by measuring SAG expression in mouse embryonic tissue (provided by Dr. Tom Glaser, University of Michigan), using reverse transcription of total RNA followed by PCR with the following primers: SAGTA.01 5'-CGGGATCCCCATGGCCGACGTGAGG-3' (SEQ ID 7) and SAGT.02 5'-CGGGATCCTCATTTGCCGATTCTTTG-3' (SEQ ID 8), which flank the entire SAG coding region. The PCR reaction mixture for 11 samples contained 55 µL of 10X buffer, 22 µL of 1.25 mM dNTP, 1.1 µL of SAGTA.01 and SAGT.02, respectively, 5.5 µL of Taq DNA polymerase, 5.5 µL of ³²P-dCTP and sterile water up to 495 µl. Into each tube which contains 5 µL of first strand cDNA reverse-transcribed for total RNA isolated from mouse embryonic tissues (Sun et al. (1997), Mol. Carcinogenesis 8:49-57), 45 µL of reaction mixture was added and PCR was performed for 25 cycles (95°C for 45 sec, 60°C for 1 min and 72°C for 2 min). A 5 µL aliquot of the PCR product was denatured and separated on a sequencing gel, which was dried and exposed to X-ray film.

SAG RNA was expressed in 9.5 day old to 19.5 day old whole mouse embryos, with a higher level of expression detected between days 9.5 and 11.5. These results suggest that SAG plays a role in embryonic development.

Example 5. Cellular localization by immunofluorescence

NIH3T3 cells (ATCC CRL 1658) were plated on coverslips in 24-well culture dishes and transfected by the calcium phosphate method according to standard techniques (Sambrook et al, 1989) with the following constructs: pcDNA3.1 (Invitrogen vector pcDNA 3 with a myc-his-tag); pcDNA3.1-SAG (human SAG cDNA subcloned into the BamHI site of pcDNA3.1, downstream from the CMV promoter and upstream and in-frame with the myc-his-tag, such that upon expression, the resulting fusion protein consists of the SAG

protein followed by the myc-his tag at the carboxy-end of SAG); or pcDNA3.1-LacZ (Invitrogen). Two days post-transfection, cells were washed once with cold PBS and then fixed with 3% formaldehyde in PBS for 10 minutes followed by 5 minutes in 1:1 methanol:acetone. The fixed cells were washed 4 times in PBS and incubated with antibody directed against the Myc-tag (Invitrogen 1:200 dilution) in PBS containing 1% BSA, 0.1% saponin, 2 μg/mL DAPI for 1 hour in the dark with shaking. Cells were then washed 4 times with 0.1% saponin in PBS and incubated with FITC-conjugated goat anti-mouse antibody (Jackson Laboratory, 1:100 dilution) for 1 hour in the same conditions as the first antibody. After incubation cells were washed 4 times with 0.1% saponin in PBS and twice with PBS. The coverslips were then mounted to glass slides with non-fade mounting medium and analyzed using a Leita Dialux 20 microscope.

SAG fusion protein was detected in both the cytoplasm and nucleus, while the β -galactosidase control was expressed predominately in the cytoplasm. No immunofluorescence staining was detected with the vector-only control. The cytoplasmic/nuclear localization of SAG was confirmed also in a SAG stable transfectant using both SAG and myc-tag antibodies. These data demonstrate that exogenously expressed SAG fusion proteins can be detected within transfected cells by using antibodies directed against an epitope fused to SAG protein.

Example 6. Expression and purification of SAG protein in bacteria

5

10

15

20

25

30

The entire open reading frame of the human SAG cDNA was PCR amplified as described above and subcloned into the pET11 expression vector (Novogen) under control of the T7 promoter, yielding construct pET11a-hSAG. The sequence and orientation of the SAG DNA insert were confirmed by DNA sequencing. pET11a-hSAG was used to transform *E. coli* strain BL21 (Novagen, Inc.). Transformed cells were grown in LB media containing ampicillin (50 µg/mL). SAG expression was induced by 0.5 mM IPTG and SAG protein was found in inclusion bodies, which were subsequently isolated as follows.

Following IPTG induction, four liters of cells were grown for 4.5 hours at 37°C at a shaker setting of 150 rpm. Cell pellets were obtained by centrifugation at 5000 rpm for 10 minutes, and were resuspended in 100 mL TN buffer (20 mM Tris-HCl, pH 7.5, 50 mM NaCl) containing 100 µM PMSF. The resuspended cell pellet was subsequently sonicated (15 sec/round for 5 rounds at a setting of 15 on Model 50 sonic dismembrator, Fisher Scientific) and subjected to pressure of 2500 pounds/square inch on a French cell press,

followed by addition of 1 mM MgCl2 and 10 mg of DNase I. The cell lysate was placed on ice for 30-60 minutes and then centrifuged at 18,000 rpm and the supernatant was disposed.

5

10

15

20

25

30

The pellet was seen to have 2 layers. The white layer on the top was carefully blown loose with TN buffer and removed. The remaining dark brown layer on the bottom was resuspended thoroughly in 15 mL of urea buffer (7 M urea, 20 mM Tris-HCl, pH 7.5, 200 mM NaCl) and allowed to sit overnight at room temperature. The resuspended cell pellet was vigorously homogneized with a serological pipette and then centrifuged at 40,000 rpm for 40 minutes using an SW50 ultracentrifuge rotor. The supernatant was collected and concentrated using a Centricon-10 concentrator to a volume of 5 mL and loaded onto a Sephacryl-100 column (100 cm long with a diameter of 2.5 cm) that had been equilibrated with urea buffer. The column was run at a rate of 0.25 mL/min and fractions were collected. The early fractions containing a brownish color consisted of mostly the large molecular weight protein, as expected. They also contained a protein with the same size of SAG protein (approximately 13 kDa). Since SAG protein contains 12 cysteine residues, it follows that SAG protein may form oligomers when expressed in bacteria and thus may elute as a SAG protein oligomer. Since SAG is a redox-sensitive protein, the DTT present in SDS sample buffer reduces SAG protein oligomers to monomer, leading to the detection of a fast migrating band. When early fractions were run in SDS-PAGE without DTT, the 13 kDa SAG protein band disappeared, and a 260 kDa band was detected, representing a SAG protein 20-mer. This unique feature helped us to purify SAG protein. Early fractions were pooled and loaded on the same Sephacryl-100 column pre-equilibrated with 7M urea and 5mM DTT.

SAG protein oligomer was reduced to monomer by using DTT in the loading buffer and was eluted in the later fractions, thus separating it from high molecular weight contaminant proteins (eluted earlier). The brownish fractions were pooled and concentrated using a Centricon-10 to a volume of 5 mL. DTT was added to a concentration of 5 mM. The combined fractions were loaded onto an S-100 column (100 cm long with a diameter of 2.5 cm), that had been equilibrated with urea buffer plus 5 mM DTT. The column was run at a rate of 0.25 mL/min and fractions were collected. The fractions containing SAG protein are brownish in color, highly suggesting that SAG is a heme-containing protein. The SAG protein containing fractions and their sensitivity to DTT were confirmed by Western blot using SAG antibody. The brownish fractions were pooled and concentrated using a Centricon-10 concentrator to a volume of 2 mL. The resulting sample was dialyzed against 4 liters of dialysis buffer (150 mM KCl, 20 mM Tris-HCl, pH 7.5) at 4°C overnight to

remove urea and DTT to yield refolded SAG protein. The dialyzed sample was loaded onto an S-100 column (100 cm long with a diameter of 2.5 cm), that had been equilibrated with dialysis buffer. The brownish fractions were pooled and concentrated using a Centricon-10 to a volume of 1 mL. The resulting sample was stored at 4°C. The protein concentration was determined by a BioRad protein assay. The purity of the samples was demonstrated in 10-20% SDS-PAGE. These data demonstrate the purification of recombinant SAG protein.

Example 7. Redox Sensitivity of SAG Protein

5

10

15

20

25

30

To confirm that purified recombinant SAG protein possesses the same redox sensitivity as it shows during protein purification, the sensitivity of refolded SAG to redox reagents was examined next. SAG protein (1 μ g) was exposed to various concentrations of DTT (1 M, 300 mM, 100 mM, or 30 mM) or H_2O_2 (15 mM, 50 mM, 150 mM or 450 mM) for 10 min before being separated by polyacrylamide gel electrophoresis (PAGE), followed by Western blot analysis. Alternatively, 10 μ g of SAG protein was incubated with 50 mM H_2O_2 for 10, 30, 60 or 120 minutes followed by PAGE separation and Coomassie Blue staining.

Dimers of SAG protein are rather resistant to reducing reagent DTT since no significant dimer was reduced to monomer after DTT treatment. However, as little as 15 mM H₂O₂ induces oligomerization of SAG protein, possibly through the formation of intermolecular disulfide bonds. The oligomerization is incubation-time dependent, as higher order SAG protein oligomers were detected upon increased incubation time. Interestingly, a band migrating faster than the monomer form is observed upon H₂O₂ treatment, and the monomer form of SAG protein becomes a doublet, possibly due to the formation of intramolecular disulfide bonds.

In order to determine whether H₂O₂-induced SAG protein oligomerization can be reversed by DTT treatment, 1 µg of purified SAG protein was incubated with 50 mM H₂O₂ for 10 minutes, followed by a 10 minute incubation with either H₂O₂, 50 mM DTT, 100 mM, 500 mM, or 1 M DTT. The samples were separated via PAGE followed by Western analysis. The results demonstrated that H₂O₂-induced SAG protein oligomerization can be reversed by subsequent incubation with DTT in a dose dependent manner, indicating that SAG protein oligomerization is subject to redox regulation.

To confirm that SAG protein oligomerization and doublet formation is due to interand intra-molecular disulfide bond formation, respectively. SAG protein was treated, prior to H₂O₂ exposure, with 50 mM N-ethylmaleimide (NEM), an alkylating reagent that will alkylate the free SH-groups in SAG protein. Purified SAG protein (1 µg) was pre-incubated with 50 mM NEM or DMSO, or buffer only, for 10 minutes prior to H2O2 treatment. The samples were separated via PAGE, followed by Western blot analysis. Pre-incubation of SAG protein with DMSO did not affect H2O2-induced oligomerization and doublet formation, whereas NEM pre-treatment abolished H2O2 activity. Neither inter-(oligomerization) nor intra- (doublet monomer) disulfide bonds were formed, demonstrating that alkylation of the free SAG protein SH groups abolishes H2O2 sensitivity. These data demonstrate that SAG protein is redox sensitive. It is subjected to both intra- and intermolecular disulfide bond formation upon exposure to H2O2, as evidenced by both doublet and oligomer formation. These H₂O₂-induced changes can be reversed by subsequent treatment with reducing reagents, including DTT, or can be prevented by NEM pretreatment. It has also been observed that zinc can promote H2O2-induced oligomerization, although zinc itself did not induce oligomerization.

Example 8: Production of SAG mutants

5

10

15

20

25

30

In order to understand the role of each particular cysteine residues in heme binding and SAG oligomerization, a series of single and double SAG mutants were made in heme binding sites as well as the zinc ring finger motif (see Figure 1B). To generate single point mutations in SAG cDNA, 15 pairs of sense and antisense primers were designed, which are partially complimentary and contain a desired point mutation. The wildtype SAG cDNA cloned into the pET11a vector at the Nhe I/Bam HI sites was used as the template for PCR amplification. Two separate PCR reactions were conducted using a) primer SAG P.01 (5'-TATGGCTAGC ATGGCCGACGTGGAGG-3) (SEQ ID 9) and each of antisense primers and b) each of sense primers and SAG T.02 (SEQ ID 8), respectively. The resultant PCR products that overlap with each other and contain a desired point mutation were mixed and served as templates for a third PCR. The primers used were SAG P.01 and SAG T.02, which flank the entire encoding region of SAG cDNA. The PCR was performed as previously described (Sun et al. (1992) BioTechniques 12:639-640). The PCR products were digested with restriction enzymes Nhe I and Bam HI and subcloned into the pET11a vector, which was digested with the same restriction enzymes. To generate SAG double mutants

(MM10, MM13, MM14, see Figure 1B), a QuickChange site-directed mutagenesis kit was purchased from Strategene (La Jolla, CA) and used as instructed. All SAG mutants generated were verified by DNA sequencing (SEQ ID 21, SEQ ID 23, SEQ ID 25, SEQ ID 27, SEQ ID 29, SEQ ID 31, SEQ ID 33, SEQ ID 35, SEQ ID 37, SEQ ID 39, SEQ ID 41, SEQ ID 43, SEQ ID 45, SEQ ID 47 and SEQ ID 49). The predicted mutant SAG proteins encoded by these mutant SAGs are shown in SEQ ID 22, SEQ ID 24, SEQ ID 26, SEQ ID 28, SEQ ID 30, SEQ ID 32, SEQ ID 34, SEQ ID 36, SEQ ID 38, SEQ ID 40, SEQ ID 42, SEQ ID 44, SEQ ID 46, SEQ ID 48, and SEQ ID 50.

5

10

15

Individual SAG mutant-expressing vectors were used to transform *E.coli* strain BL21 (Novagen, Inc.). Mutant SAG protein was expressed and purified as detailed in Example 6. The fractions after a Sephacryl-100 column were collected and analyzed on 8-25% Phast gels followed by Coomassie blue protein staining. The pure fraction containing mutant SAG protein was dialyzed in 4 liters of 20 mM Tris-HCl, pH 7.5 and used for SAG protein oligomerization studies.

Purified wildtype SAG protein is a heme-containing brownish protein (See Example 9). Some of the purified SAG protein mutants were found to have either lost the brownish color (MM3 and MM13) or had decreased brownish color (MM1) compared to wildtype SAG protein. This color change indicates the loss or decrease of heme binding (Table 1).

TABLE 1. SUMMARY OF SAG MUTANTS

NAME	MUTATION SITE(S)	HEME BINDING	OLIGOMERIZATION
WT	None	+++	Yes
MM1	C _A /heme	++	Yes
MM2	C _B /heme	+++	Yes
MM3	C _{A+B} /heme	+/-	Yes
MM4	C ₁ /Zn-ring finger 1	+++	Yes
MM5	C ₃ /Zn-ring finger 1	+++	Yes
MM6	H ₄ /Zn-ring finger 1	+++	Yes
MM7	H ₅ /Zn-ring finger 2	+++	Yes
MM8	C ₆ /Zn-ring finger 2	+++	Yes
ММ9	C ₇ /Zn-ring finger 2	+++	Yes
MM10	H ₄₊₅ /Zn-ring fingers 1&2	+++	Yes
MM11	C ₂ /Zn-ring finger 1	+++	Yes
MM12	C _c /protease inhibitor	+++	Yes
MM13	C ₁₊₂ /Zn-ring finger 1	+/-	Yes
MM14	C ₇₊₈ /Zn-ring finger 2	+++	No
MM15	GADPH binding site	+++	Yes

To examine mutant SAG protein oligomerization, each mutant SAG protein as well as wildtype SAG was treated with 50 mM H₂O₂ for 10 min. All of the SAG mutants, except MM14, can be oligomerized upon exposure to H₂O₂. The mutant 14, which is a double mutants in positions of C7 and C8 in the zinc ring finger domain, becomes insensitive to oligomerization (Table 1), indicating that these two positions are important for intermolecular disulfide bond formation.

Example 9. Heme measurement of SAG protein

5

10

Heme content in SAG protein was measured as previously described (Rieske (1967) Methods in Enzymol. 76, 488-493). Briefly, 1 mg of purified SAG protein, along with cytochrome C, catalase, and BSA as controls, was extracted with cold acetone (0.5 mLs)

After centrifugation the pellet was extracted sequentially with 0.5 mL of chloroform:methanol (2:1); 0.5 mL of cold acetone, and finally 0.5 mL of cold acetone containing 5 µL of 2.4 N HCl. The acetone extracts were dried under speed-vac and dissolved in 0.5 mL of pyridine. After addition of 0.5 mL of 0.2 N NaOH, the solution was centrifuged briefly and clear supernatant was recovered. One drop of diluted potassium ferricyanide (0.05 M) was added to the supernatant and the absorbance was read at 556 nm in 1.0 mL quartz cuvettes using water as a blank. The solution was then reduced by adding 10 uL of 2 M DTT and absorbance was read at 556 nm, 587 nm and 550 nm, respectively.

Heme absorbance at 556, 587, and 550 nm was observed in SAG protein, as well as in cytochrome C and catalase, but not in BSA. This result demonstrated that SAG protein contains heme, but did not reveal the molar ratio between SAG protein and heme molecule.

Example 10. SAG protein antibody production

5

10

15

20

30

Two polyclonal antibodies against SAG protein were generated using standard methods [by Zymed Laboratories, Inc. (San Francisco) under a service agreement with Warner-Lambert]. Briefly, the peptide antibody was generated as following. A 16-amino-acid peptide (SAG-Pep1: QNNRCPLCQQDWVVQR) (SEQ ID 10) located in the C terminus of SAG protein (codons 95-110) was synthesized and purified via standard techniques. The purified peptide was conjugated to keyhole limpet hemocyanin (KLH) via cysteine residues. The conjugated peptide (0.5 mg) was emulsified with equal volume of Complete Freund Adjuvant (CFA) and subcutaneously injected into rabbit, followed by 4 boosts with 0.5 mg each in Incomplete Freund Adjuvant (IFA) at 3 week intervals. Rabbits were bled 10 days after the final boost and antiserum was collected. The same protocol was used for protein antibody production using purified human SAG protein as the antigen, prepared as described above.

25 Example 11. Analysis of SAG protein transcriptional regulatory activity

SAG protein belongs to the zinc ring finger protein families by virtue of its C₃H₂C₃ motif (Saurin et al. (1996) TIBS 21, 208-214). Some zinc ring finger proteins have been shown to bind to DNA and function as transcriptional repressors (for example, RING1) (Satijn et al. (1997) Mol. Cell. Biol. 17, 4105-4113), whereas others function as transcriptional activators (Chapman and Verma (1996) Nature 382, 678-679; Monteiro et al. (1996) Proc. Natl. Acad. Sci. USA 93, 13595-13599). To examine the transcriptional regulatory activity of SAG protein, the cDNA encoding the entire open reading frame of

human SAG was PCR amplified and fused both in frame and as an antisense fusion, downstream of the Gal-4 DNA binding domain (encoding amino acids 1-147) in the pG4 vector (Sadowski et al., Nature 335:563-564, 1988). The resulting construct was sequenced to confirm in frame fusion and freedom from PCR-generated mutation. The construct was co-transfected along with a chloramphenicol acetyltransferase (CAT)-reporter-expressing vector (Sadowski et al., Nature 335:563-564, 1988) as well as a β-galactosidase reporter whose expression is driven by a CMV promoter for normalization of transfection efficiency into human kidney 293 cells (ATCC accession number CRL1573) by the calcium phosphate method. CAT activity was measured 36 hours post-transfection using a CAT assay kit (Quan-T-CAT; Amersham) according to the manufacturer's instructions. PG4-VP16, a known transcription factor (Triezenberg et al., Genes and Develop. 2:718-729, 1988), fused downstream of the Gal4 DNA binding domain was used as a positive control. Activation was calculated by arbitrarily choosing CAT activity from the vector control as 1 and comparing the other constructs to it. Three independent transfections and assays were performed.

SAG protein showed no transactivation activity. The positive control, VP16 showed 300-fold activation of CAT activity. To test for transrepression activity, SAG constructs (both sense and antisense) were co-transfected with pG4-VP16. Again, neither orientation of SAG induced significant expression of VP16-induced transactivation. These results demonstrated that SAG protein lacks transcriptional regulatory activity when fused downstream Gal-4 DNA binding domain.

Example 12. SAG is an RNA binding protein

5

10

15

20

25

30

The zinc-ring finger domain of the MDM2 protein has been shown to bind to RNA (Elenbaas et al. (1996) Mol. Med. 2, 439-445). Since SAG protein showed no transcriptional regulatory activity, it was tested whether SAG protein could bind to RNA or DNA. Binding of purified SAG protein to different nucleic acid cellulose conjugates was performed as described (Elenbaas et al. (1996)). Briefly, 0.5 µg of SAG protein was incubated in 300 µL RNA binding buffer for 1 hour at 4°C with double-stranded calf thymus DNA, denatured calf thymus DNA (ssDNA), or one of 4 RNA homopolymer columns (Sigma) conjugated to agarose or cellulose beads (Sigma), and used according to the manufacturer's instructions. RNA binding buffer consisted of 20 mM Tris, pH 7.5, 150 mM NaCl, 5 mM MgCl₂, 0.1% nonidet P-40, 50 µM ZnCl₂, 2% glycerol, and 1 mM DTT. The columns were washed with 3 mL RNA binding buffer to remove non-specifically bound protein from the beads, which

were then boiled in SDS sample buffer. The protein so eluted from the beads was separated by SDS-PAGE, transferred to nitrocellulose for Western blot analysis using the polyclonal antibody directed against SAG protein described previously detected by ECL chemiluminescence (Amersham) according to the manufacturer's instructions.

Purified SAG bound to polyU, polyA, and polyC RNA, respectively. No binding was seen with polyG RNA or ssDNA. A band showing dsDNA binding did not agree with SAG molecular weigh. Oligomeric SAG protein bound to polyU RNA, whereas the monomeric form of SAG binds to polyA and polyC RNA. Purified SAG protein was run as a marker. These results suggest that SAG is an RNA binding protein and that binding specificity is determined by the oligomeric form of SAG protein.

Example 13. Identification of two deletion mutants of SAG in cancer cell lines

5

10

15

20

25

30

Total RNA was isolated from DLD-1 colon carcinoma cells (ATCC accession number CCL221) and subjected to RT-PCR using primers SAG TA.01 and SAG T.02. The resulting PCR fragments were subcloned into the TA cloning vector (Invitrogen). During sequence verification of the resulting clones, it was found that several clones contained either a 7 bp or a 48 bp deletion at nucleotide 170 or 177, respectively, assigning the first A at the start codon as nucleotide #1. Both SAG deletions encode the potential heme-binding sites. The 7 base pair deletion (SAG mutant 1) (SEQ ID 11) is a frame shift deletion that abolishes the downstream encoded zinc-ring finger motif in the resulting protein (SEQ ID 12), whereas the 48 base pair deletion (SAG mutant 2) (SEQ ID 13) is an in-frame deletion that eliminates 16 amino acids in the encoded protein (SEQ ID 14), but retains the zinc-ring finger motif.

Total RNA was isolated from a total of 20 human tumor lines and transformed lines originating from lung, brain, kidney, prostate, testis, nasopharynx, bone, cervix and foreskin and subjected to RT-PCR analysis as described previously (Sun et al. (1993) Mol. Carcinogenesis 8, 49-57). Genomic DNA was also isolated from these cell lines and subjected to PCR amplification as described (Sun et al. (1992) BioTechniques 12:639-640). The primers used for PCR were hSAG.M1, 5' GCCATCTGCAGGGTCCAG-3' SAGT.02-1 151 of hSAG cDNA. and (SEO ID 15), starting at nt 5'-GGATCCTCATTTGCCGATTCTTTGGAC-3' (SEQ ID 16), including stop codon (underlined). The resulting fragment is 200 bp for wildtype SAG. The PCR was conducted in the presence of 35S-dATP (Amersham) and PCR products were resolved in 6% denaturing sequencing gels, as described previously (Sun et al. (1995) Cancer Epidemiology, Biomarkers & Prevention, 4, 261-267). The bands corresponding to wildtype as well as the

DNIS no-

two deletion mutants were cut out from the gel, PCR amplified using the same set of primers, and sequenced to verify the DNA sequence of the resulting PCR fragments.

Both the 7 base pair and the 48 base pair deletions were detected in RNA from only the CATES-1B cell line, a testicular carcinoma line obtained from ATCC (accession number HTB104). This tumor line also contains the wildtype SAG DNA sequence. The identity of these three bands was confirmed by DNA sequencing after PCR amplification and TA cloning. HONE-1, a nasopharyngeal carcinoma line which only contains wildtype SAG was included for comparison.

It was next examined whether these SAG deletions were detectable at the DNA level. Genomic DNA was isolated from CATES-1B cells and subjected to PCR analysis, as described previously (Sun et al. (1992) BioTechniques 12:639-640). The primers used were hSAG.M1 and SAG T.02 (see above for sequences). Genomic DNA from CATES-1B cells possesses only wildtype SAG and no SAG deletion mutants were detected. These results indicate that the SAG deletion mutations occur very rarely in human cancer lines. Detection of the mutations in SAG RNA, but not genomic DNA, may reflect an RNA editing modification of SAG messenger RNA.

Example 14. Production of stable SAG transfected mammalian cells

5

10

15

20

25

30

The potential biological function of human SAG protein was examined next by its overexpression in cells. DLD-1 cells were transfected with the following plasmids: the neo control pcDNA-3 (Invitrogen) (identical to pcDNA3.1 described above, except that it lacks the myc-his tag), pcDNA-SAG, pcDNA-SAG-mutant-1, and pcDNA-SAG-mutant-2 (pcDNA3 with SAG, SAG 1 or SAG 2 subcloned into the BamHI site, respectively, using methods well known in the art). The SAG mutant constructs were generated by RT-PCR as follows. Total RNA was isolated from DLD-1 cells, and subjected to reverse transcription, followed by PCR amplification. The primers used were SAG.TA01 (SEQ ID 7) and SAGT.02 (SEQ ID 8), which flank the entire coding region of SAG gene. The PCR products were digested with restriction enzyme Bam HI, and subcloned into pcDNA3 (In Vitrogen, San Diego), a mammalian expression vector under the transcriptional control of the CMV promoter, which drives gene expression constitutively. The resultant clones were sequenced to confirm both sense and antisense orientation and freedom of PCR-generated mutations. DNA sequencing revealed wildtype SAG clone as well as two deletion mutants: SAGmutant-1 (7 bp deletion, SEQ ID 11) and SAG-mutant-2 (48 bp deletion, SEQ ID 13) in DLD-1 tumor cells.

DLD-1 cells were transfected by lipofectamine (BRL) with plasmids expressing wildtype (both sense and antisense orientation), SAG mutant-1, and SAG mutant-2, along with the neo control vector. Neomycin resistant colonies were identified by G418 selection (600 µg/mL) for 18 days. Stable clones were ring-isolated by well known methods (Sun et al. (1993) Proc. Natl. Acad. Sci. USA. 90: 2827-2831) and SAG expression was monitored by Northern analysis. Selected clones were examined for SAG protein expression by immunoprecipitation, as described below.

5

10

15

20

25

Total RNA was isolated from the cloned cell lines and subjected to Northern analysis. Cell lines transfected with the following constructs were analyzed: vector controls D1-3 and D1-6; SAG-wildtype D12-1 and D12-8; SAG-mutant-1 D3-3 and D3-4; and SAG-mutant-2 D4-2 and D4-5.

Northern blot analysis of RNA from selected stable SAG-expressing clones probed with the human SAG cDNA demonstrated that all SAG transfectants express SAG mRNA, while very low levels of endogenous SAG message were detected in the neo control cells.

The vector control lines and SAG wildtype and SAG deletion mutant transfectants were subsequently subjected to immunoprecipitation using standard techniques (Sun et al. (1993) Proc. Natl. Acad. Sci. USA. 90: 2827-2831, Sun et al. (1993) Mol. Carcinogenesis 8, 49-57). Subconfluent SAG transfectants were subjected to methionine starvation for 1 hour and then metabolically labeled with ³⁵S-translabel (0.2 mCi/mL) for 3 hours. Cells were then lysed on ice for 30 minutes in a lysis buffer comprising 2% Nonidet P40, 0.2% SDS, 0.5% sodium deoxycholate, 1 mM sodium orthovanadate, 5 mM sodium fluoride, 5 mM sodium pyrophosphate, 1 mM phenylmethylsulfonyl fluoride, and 1 µl/mL leupeptin, and centrifuged at 12,000 x g. The TCA precipitable radioactivity in the supernatant (1 x 108 cpm) was immunoprecipitated using rabbit anti-human SAG antibody (generated as described above). The immunoprecipitates were collected, washed, and analyzed on a 10-20% SDS-polyacrylamide gel, followed by autoradiography. High SAG protein expression was detected only in the wildtype transfectants. The antibody used did not recognize the two SAG protein mutants. These data demonstrate the production of stably transfected cells expressing either wildtype or mutant SAG protein

30 Example 15. Morphological appearance of SAG transfectants after exposure to redox reagents

Two neo controls (D1-3 and D1-6) and two SAG-producing lines (D12-1 and D12-8) were chosen to examine their sensitivity to redox compounds by morphological observation. After exposure to 150 μM OP, 200 μM H₂O₂, or 125 μM zinc for 24 hours, the neo-control cells were shrunken and detached, a sign of apoptosis, while SAG-expressing cells appeared morphologically normal. These results indicate that SAG production protects cells from apoptosis induced by redox compounds. Expression of SAG, however, did not offer the protection against copper. No difference in morphological signs of apoptosis was observed with CuSO₄ treatment (up to 750 μM) between the vector controls and SAG transfectants. Higher doses induced apoptosis in all lines.

Example 16. SAG expression protects cells from DNA fragmentation

5

10

15

20

25

30

The sensitivity of these SAG-transfected cells to OP-induced apoptosis was examined next by monitoring DNA fragmentation, a hallmark of apoptosis. Subconfluent (80-90%) SAG transfected cells expressing wildtype SAG, SAG mutant-1, SAG mutant-2, or vector control cells, were seeded at 3.5 x 10⁶ per 100 mm dish and exposed after 16-24 hours to 150 μM OP, 125 μM zinc sulfate, or 200 μM H₂O₂ for 24 hours. Both detached and attached cells in 2 x 100 mm dishes were harvested and subjected to DNA fragmentation analysis as follows. Cells were collected by centrifugation and lysed with lysis buffer (5 mM Tris-HCL, pH 8; 20 mM EDTA; 0.5% Triton-X100) on ice for 45 minutes. Fragmented DNA in the supernatant of a 14,000 rpm centrifugation (45 minutes at 4°C) was extracted twice with phenol/chloroform and once with chloroform and precipitated by ethanol and salt. The DNA pellet was washed once with 70% ethanol and resuspended in TE buffer with 100 μg/mL RNase at 37°C for 2 hours. The fragmented DNA was separated in 1.8% agarose gel electrophoresis, stained with ethidium bromide, and visualized under ultraviolet light.

OP induced apoptosis in the vector control cells. Less DNA fragmentation was observed in wild type SAG transfected cells compared to control cells. SAG mutant 1, which does not encode the zinc ring-finger motif, did not show any protection against OP-induced DNA fragmentation, whereas SAG mutant 2, which retains the zinc ring finger domain, still showed protection. These results suggest that overexpression of SAG protein protects cells against OP-induced apoptosis, and the zinc ring finger domain is required for this protective activity.

Since SAG protein contains a zinc ring finger motif, the sensitivity of SAG transfectants to zinc treatment was examined next. Zinc induced apoptosis in DLD-1 cells

transfected with the vector only. Induction of apoptosis was limited by SAG overexpression, which showed much less DNA fragmentation than the control lines. This data suggests that the SAG protein binds to and chelates zinc through the zinc ring finger domain and thus provides increased resistance to zinc toxicity compared to non-transfected cells.

Another feature of SAG is the formation of oligomers after exposure to H_2O_2 . Cells may be protected from H_2O_2 induced toxicity by SAG oligomerization. SAG-transfected cells were, therefore, treated with H_2O_2 followed by assays for DNA fragmentation. H_2O_2 induced apoptosis in DLD-1 cells. SAG protein overexpression partially protected cells from H_2O_2 -induced apoptosis, as evidenced by a reduction in DNA fragmentation. Taken together, these results demonstrate that SAG affords at least some protection against apoptosis induced by redox compounds such as OP and H_2O_2 and also against apoptosis caused by zinc.

Example 17. Antisense SAG expression inhibits tumor cell growth

5

10

15

20

25

30

To test the growth effects induced by SAG expression, DLD-1 cells were transfected with the neo control vector, or vectors expressing SAG, SAG mutants 1 or 2, or antisense SAG, as described above. Neomycin resistant colonies were selected with G418 (600 µg/mL) for 18 days and stained with 50% methanol/10% acetic acid/0.25% Coomassie Blue.

A stable DLD-1 transfectant expressing antisense SAG mRNA (D15-1) was cloned after G418 selection in order to examine potential changes in tumor cell phenotype caused by decreased SAG expression. Subconfluent D15-1 cells, along with the vector control cell (D1-6), and SAG (sense) overexpressing cells (D12-1 and D12-8) were metabolically labeled and subjected to immunoprecipitation using SAG protein antibody as described above. Densitometric quantitation of SAG protein expression using a computing densitometer, (Molecular Dynamics) was performed according to the manufacturer's instructions. The number was calculated by arbitrarily choosing the value from the vector control cell D1-6 as 1. Antisense SAG transfected cells (D15-1) exhibited a 60% reduction in endogenous SAG protein. Monolayer growth of DLD-1 cells was significantly inhibited by antisense SAG transfection. None of the other transfectants were growth-inhibited, as compared to the neo control.

It was next examined whether antisense SAG-transfected cells would exhibit growth inhibition in soft agar. D15-1 cells, along with transfectants expressing wildtype SAG (D12-8), SAG mutant-1 (D3-3), SAG mutant-2 (D4-2), as well as the neo control (D1-3)

were grown in 0.25% agar medium for 14 days. Colonies containing greater than 16 cells were counted. Three independent experiments, each run in duplicate, were performed. Shown is the mean +/- standard error of the mean. As shown in Figure 2, down-regulation of SAG in D15-1 cells did cause significant growth inhibition of DLD-1 cells as reflected by 75% reduction of soft agar colony number when compared to the neo control (D1-3), SAG (sense) expressing line, D12-8, and SAG mutants (D3-3, D4-2).

5

10

15

20

25

30

In a further study, 4 x 10⁶ confluent D15-1 cells along with parental DLD-1 cells, the vector control D1-6, and SAG wildtype transfectant D12-1 cells were inoculated subcutaneously into SCID mice (Taconic Farms, Germantown, New York), 10 mice per group. Tumor growth was observed twice a week. The average tumor size/mass for 10 mice was plotted against time post injection up to 24 days. When implanted into SCID mice, antisense expressing line D15-1 failed to form tumors up to 24 days after inoculation, whereas substantial tumor growth was observed in parental DLD-1 cells, the neo control D1-6 cells, and SAG (sense) expressing D12-1 cells (Figure 3). All these experiments demonstrate that downregulation of SAG expression leads to growth inhibition of tumor cells, and further indicates that SAG is a cellular protective molecule.

Example 18. Cancer gene therapy using adenovirus expressing antisense SAG

Since antisense SAG expression has been shown to inhibit tumor growth both *in vitro* and *in vivo* (example 17), SAG can be used as a target for cancer gene therapy. Methods for conducting cancer gene therapy are well known in the art (see Zhang and Fang, Exp. Opin, Invest. Drugs 4: 487-514, 1995 and Zhang et al., Adv. Pharmacol. 32: 289-341, 1995).

Tumor cell lines with endogenous SAG expression, including, but not limited to DLD-1 (colon), Du145 (prostate), G401 (kidney), H2009 (lung) and HONET-1 (nasopharynx), are used to establish the tumor models,. Tumor cells from tissue culture are suspended in PBS at a concentration of 5 x 10⁷/mL and stored on ice. 0.2 mL of the cell suspension (containing approximately ten million cells) is subcutaneously injected into the flank of 6- to 8-week-old athymic nude mice and tumors are allowed to grow for 30-40 days or until the average tumor size reaches 5 mm.

Recombinant adenoviral vectors expressing antisense human SAG, driven by the CMV promoter (Ad.CMV-SAG) were produced by co-transfecting a shuttle plasmid (pJM17, circularized Ad5 genome) and a recombinant plasmid (pEC-SAG; a CMV driven plasmid containing left arm of Ad5 genome) into 293 cells.

31

Tumors are injected with either 0.1 mL of recombinant adenoviral solution (1-5 x 10¹⁰ pfu/mL) or 0.1 mL of PBS alone as a control. Daily treatment is performed for 2 days and after 1 week without treatment, daily treatment is resumed for 3 days. The tumor size is measured daily for 2 weeks. To test combinatorial therapy with oxygen radical-generating reagents or irradiation, the treated group is subdivided into three sub-groups (10 mice per subgroup): group A receives adenovirus alone (see above); group B receives adenovirus and at the same time receives an intraperitoneal injection of adriamycin (3 mg/kg) an oxygen radical-generating reagent, and group C receives adenovirus plus irradiation at 350cGy of cesium-137. Some tumor-bearing mice will only receive the same dose of adriamycin or irradiation as drug or irradiation controls,.

5

10

15

20

25

30

Expression of antisense SAG blocks endogenous SAG synthesis, which renders tumor cells supersensitive to oxygen radicals. Significant tumor shrinkage in treated tumors with or without drugs or radiation, as compared with the vehicle control, indicates the efficacy of this therapy. The tumors in both control and treated groups can be further examined histologically. Samples can be immediately embedded in optimal cutting temperature compound (Miles, Inc. Elkhart, Indiana) and snap-frozen in liquid nitrogen for frozen section preparation (3-5 μm) for enzymatic staining (e.g., terminal deoxynucleotidyl transferase (Boehringer Manheim, Indianapolis, Indiana) staining for apoptosis) or immunohistochemical staining for expression of the antisense SAG. Alternatively, the samples may be fixed in 10% formalin for histologic sectioning and analyze with hematoxylin-eosin (Sigma, St. Louis, Missouri) staining.

Example 19. SAG functions as a oxygen radical scavenger to prevent oxygen radical induced damages

SAG protein contains 12 cysteine residues and forms disulfide bonds both intermolecularly and intramolecularly after exposure to hydrogen peroxide. SAG protein also binds to heme, which can modulate oxidants by oxidation/reduction of Fe(++). This oxidative buffering activity may qualify SAG as an oxygen radical scavenger.

Yeast cells having deletions in antioxidant enzyme genes [superoxide dismutase (SOD) and catalase (CAT)] are supersensitive to superoxide anion and hydrogen peroxide (Longo et al. (1997), J. Cell Biol. 137:1581-1588). Yeast cells that lack (a) Cu, Zn-SOD, (b) Mn-SOD, (c) both Cu, Zn-SOD and Mn-SOD, and (d) CAT have been transfected with human SAG expression plasmids. Sensitivity of these transfected cells to oxygen radical producing compounds such as paraquat (a superoxide anion generating compound) and

hydrogen peroxide are tested in yeast growth assays and compared to the growth of the same host cells transfected with vector controls. Rescue of these yeast cells from oxygen radical-induced cell killing indicates that SAG is an effective oxygen radical scavenger.

Example 20. Prevention of IL-1\beta induced brain injury during ischemia by SAG administration

5

10

15

20

25

30

It has been previously shown that middle cerebral artery occlusion in rats causes overexpression of interleukin-1 which induces brain injury by the release of free radicals (Yang et al., Brain Research 751:181-188, (1997)). Two experiments are conducted to test whether SAG, by scavenging free radicals released, will prevent brain damage.

In the first experiment, human SAG is subcloned into an adenovirus vector driven by RSV promoter (AdRSV-SAG). The adenoviral suspension is injected stereotactically into the lateral ventricle to ensure SAG expression in brain. Five days after administration of adenovirus, middle cerebral artery is occluded in animals for 24 hours as described (Yang et al., Brain Research 751:181-188, (1997)). Brain edema (as measured by brain water content) and cerebral infarct size, measured by histological techniques (Yang et al., Stroke 23:1331-1336, (1992)) is determined. As compared to the vector control, any reduction of brain edema and infarction size indicates SAG protection against free radical induced damage.

In the second experiment, middle cerebral artery occlusion is performed with the rat suture model, allowing either permanent (6 hours) or temporary occlusion (3 hours of occlusion and 3 hours of reperfusion) (Yang and Betz, Stroke, 25:1658-1665, (1994)). Rats then receive an injection of purified SAG protein at the size of occlusion. Brain water, ion contents, and infarct volume are measured to determine brain infarction and blood-brain barrier disruption. As compared to injection of the vehicle control, reduction in brain infarction size and blood-brain barrier disruption indicates a SAG protective effect.

Example 21. Human cancer diagnosis using SAG as a marker:

Two SAG deletion mutants in human cancer cell lines originating from colon and testis have been identitifed. Twelve pairs of colon carcinomas and adjacent normal tissues were collected from 12 patients. Genomic DNA and total RNA are isolated from these samples and subjected to PCR amplification. The resulting amplification products are analyzed for detection of SAG deletion mutations by methods well known in the art, including but not limited to RNA protection assays, DNA sequencing, hybridization, and gel

33

electrophoresis for deletion mutants. Mutations detected in tumor tissues but not in normal adjacent tissues indicate that they are tumor specific mutations and can be used as a diagnostic tool in the clinic for colon as well as testicular carcinomas.

Example 22: The yeast homolog of human SAG gene is essential for yeast growth

5

10

15

20

25

30

To further understand the function of SAG, yeast SAG knock-out mutants were constructed by homologous recombination. The construct used to knockout yeast SAG was made by PCR of a kanamycin cassette from kanMX4 plasmid (Wach et al., Yeast 10:1793-1808, 1994). The primers used for PCR were SAGKanMX4-5: 5'-TTCTCCAGTGGCAGAGAACTTTAAAGAGAAATAGTTCAAC

CGTACGCTGCAGGTCGAC-3' (SEQ ID 17), and SAGKanMX4-3: <u>5'-ACCTCGGTA</u> TGATTTAAATGTTTACGGGCAATTCATTTTT

ATCGATGAATTCGAGCTCG-3' (SEQ ID 18). The primer SAGKanMX4-5 consists of yeast SAG DNA sequence (ATCC Accession number Z74876) immediately upstream of the initiation codon ATG (underlined) and the upstream kanamycin cassette sequence at its 3'-end. Primer SAGKanMX4-3 consists of yeast SAG DNA sequence immediately downstream of the stop codon TGA (underlined) and the downstream kanamycin cassette sequence at its 3'-end.

PCR was conducted for 5 cycles at 94°C 1 min, 50°C, 1.5 min, 72°C 2 min, followed by 25 cycles at 94°C, 1 min, 56°C, 1.5 min, 72°C 2 min, followed by a 10 min extension at 72°C. The resulting PCR product (1.5 kb) was gel-purified using Qiaex II gel-purification kit (Qiagen) according to the manufacturer's instruction, and was used to transfect the diploid yeast strain Y21 using the YEASTMAKER yeast Transformation System (ClonTech Laboratory, Inc.) according to the manufacturer's instruction. Following transfection, yeast cells were grown in YPD media (Difco) containing G418 (200 μg/mL, BRL) to select transfectants containing the kanamycin cassette, which have had the yeast SAG deleted by homologous recombination.

Several G418-resistant clones were selected and assayed to determine whether heterozygous or homozygous deletions had been produced. The primers used are SAGPCR-5: 5'-TTCTCCAGTGGCAGAGAAC-3' (SEQ ID 19) and SAGPCR-3: 5'-ATGATTTAAATGTTTACGGGC-3' (SEQ ID 20). These primers constitute fragments of SAGKanMX4-5 and SAGKanMX4-3, respectively, and flank the entire yeast SAG coding region. PCR of wildtype yeast SAG produces a 0.35 kb band, whereas PCR of SAG deletion

mutants give rise to 1.5kb band, consisting of the kanamycin cassette. Both the 0.35 kb and 1.5 kb fragments were generated in all of the clones tested, indicating that heterozygous mutants were produced. Identical knock-out experiments were conducted with haploid yeast cells (InvSC1 from In Vitrogen) and no G418-resistant clone was isolated.

5

10

15

20

25

30

The failure to isolate homozygous yeast SAG deletion mutants suggests that yeast SAG is essential for growth. To confirm this, 12 individual heterozygous yeast strains (y21ySAG/ySAG::Kan) were sporulated to determine if yeast SAG-kan haploids were viable. The strains were inoculated into minimal potassium acetate sporulation media, supplemented with uracil, lysine, adenine and tryptophan (Kassir, and Simchen, G. Method Enzymol. 194, 94-110, 1991) and grown at 30°C for 7 days. Tetrads was dissected into 4 haploid offspring from each strain. For dissection, a clamp of cells from the sporulation plate was suspended in 100 µL of 1 M glycerol containing 0.5 mg/mL zymolase T20. After 30 min at 37°C, the suspension was diluted with 800 µL sterile water and put on ice. A loop of suspension was struck across a YPD plate and examined under a Zeiss Tetrad microscope for tetrads. The glass microneedle of the scope was used to dissect 4 tetrads from each strain. Two of these four haploid cell should contain wildtype SAG, while the other two should contain a yeast SAG deletion. In all 12 clones, only two out of four dissected cell grew, and none were viable in YPD medium supplemented with G418, indicating that viable cells did not contain the kanamycin cassette or the SAG deletion. The experiment clearly demonstrate that SAG is essential for yeast growth, further demonstrating its evolutionary importance.

To determine if ySAG is required for normal growth or simply for germination, hSAG was cloned into a yeast expression vector with URA3 selectable marker. The hSAG-URA plasmid was then transformed into heterozygous ySAG knockout cells, and transformants were selected on URA-minus plates. Clones expressing hSAG (measured by Western blot analysis) were sporulated and tetrads were dissected. Viable colonies were then screened on either YPD alone, or YPD+G418, or YPD+5-fluoroorotic acid (5-FOA; used to select against the URA3-containing centromere plasmid (Boeke et al., Mol. Gen. Genet, 1984;197:345). Again the hSAG-URA3 plasmid complemented the ySAG::kan allele, as all four haploids from four individual tetrads grew. When grown on YPD+G418 plates, two haploids from each tetrad die, indicating that they contain the wildtype ySAG gene. Other two haploids from each tetrads survived, indicating they contained ySAG::kan allele. When these latter colonies were grown on YPD+5-FOA plates, which selects against URA3 plasmid, all failed

to grow, indicating that ySAG is essential for normal vegetative growth and not simply for sporulation.

Example 23: Human SAG rescue of yeast SAG knockout phenotype

To examine whether human SAG can rescue death phenotype of yeast SAG knockout, wildtype human SAG, along with the SAG mutants (MM3, sequence ID 25; MM10, sequence ID 39; and MM14, sequence ID 47, Figure 1A) were constructed into a plasmid with selection marker and transfected into heterozygous yeast strain Trp (y21-SAG/ySAG::Kan) as described above. The clones grown in Trp-minus/G418-plus plates were examined by Western blot analysis for SAG expression. The clones expressing human SAG were sporulated and dissected. In 10 wildtype human SAG clones, 3 or 4 haploids are viable. Some of them contain yeast SAG, whereas the others contain ySAG K/O plus human SAG, indicating human wildtype SAG can complement yeast SAG knockout. All three mutant clones (total of 41 tested) gave rise to 1 or 2 haploids and all survival haploids contains yeast SAG, indicating that human SAG mutants cannot complement yeast SAG knockout.

Example 24: SAG binds to metals

5

10

15

20

25

30

Since SAG contains a zinc-ring finger domain, it has the potential to bind with metals. To measure potential metal binding of SAG, electrospray ionization mass spectrometry (ESI-MS) (Fenn et al., 1989) was used to compare the molecular mass of SAG under denaturing and non-denaturing solution conditions (Loo, 1997; Witkowska et al., 1995).

ESI-MS was performed with a double focusing hybrid mass spectrometer (Finnigan MAT 900Q, Bremen, Germany) with a mass-to-charge (m/z) range of 10,000 at 5 kV full acceleration potential. A position-and-time-resolved-ion-counting (PATRIC) scanning array detector was used. An ESI interface based on a heated metal capillary inlet and a low flow micro-EsI source (150 nL/min analyte flowrate) were used (Sannes-Lowery et al., 1997). The metal capillary temperature was maintained around 150-200°C for metal-protein complex studies. Recombinant protein under 7 M urea-denaturing solution was refolded by dialyzing in 50 µM ZnCl₂ for 3 days with three changes of buffer. Prior to ESI-MS measurement, the SAG solution was washed with a solution of 10 mM ammonium bicarbonate (pH 7) and 1 mM DTT, and excess zinc was removed by centrifugal ultrafiltration by passing through a 10 kDa molecular weight cut-off centrifugal filtration cartridge (Microcon-10 microconcentrator, Amicon, Beverly, MA). For the ESI-MS

analysis, a small portion of the filtered SAG protein solution was diluted into either a denaturing solvent (80:15:5 acetonitrile:water:acetic acid v/v/v, pH 2.5) or a non-denaturing solution (10 mM ammonium bicarbonate and 1 mM DTT, pH 7).

Zinc binding of SAG was first measured. Under a denaturing acidic solution (pH 2.5 and high organic concentration) where the protein is not expected to retain metal-binding characteristics even in the presence of zinc, the molecular mass of SAG was measured to be 12550, in close agreement with the expected mass for the apo-protein (12552 Da). The ESI-MS analysis of the SAG protein in a non-denaturing aqueous solution (pH 7) resulted in an increase in mass to 12733 and 12800 Da. These masses are consistent for the holo-protein binding 3 and 4 zinc metal ions, respectively.

5

10

15

20

25

30

Copper binding to SAG was also measured. As little as 1 µM CuSO₄ in the dialysis solution causes SAG precipitation with a blue (copper) color, suggesting a copper binding. Next, using ESI-MS, the potential copper binding of SAG was measured in a non-denaturing solution described above. Addition of copper acetate to a final concentration of 10 µM resulted in a further increase in mass to approximately 12929 Da. However, a precise mass could not be obtained, as a wide distribution of copper adducts appears to bind to SAG protein. Adding copper to higher concentrations resulted in precipitation of the protein.

Example 25: SAG minimizes or prevents LDL oxidation induced by copper ion or a free radical generator

Due to its H₂O₂ buffering and metal binding, it was reasoned that SAG may prevent oxidation of macromolecules induced by metal or free radical generator. An LDL (low density lipoprotein) oxidation induced by copper ion or a free radical generator, AAPH (2,2-azobis-2-amidinopropane hydrochloride), was used as a model to test potential protection activity of SAG against lipid peroxidation.

Lipoproteins (100 μg of protein/mL, Intraocel) were incubated with 10 μM CuSO₄ or with 5 mM AAPH for 4 hours at 37°C in the presence of various concentrations of purified SAG protein. AAPH is a water-soluble azo compound that thermally decomposes and generates water soluble peroxyl radicals at a constant rate (Frei et al., 1988). Oxidation was terminated by the addition of 10 μM butylated hydrozytoluence (BHT) and refrigeration at 4°C. The extent of lipoprotein oxidation was measured by the TBARS assay, using malondialdehyde (MDA) for the standard curve, as described (Buege & Aust, 1978).

5

10

15

20

25

30

Copper-induced LDL oxidation, as measured by the formation of thio barbituric acid reactive substances (TBARS), was slightly enhanced by SAG at low concentrations. At higher SAG concentrations, however, a dose-dependent inhibition (up to 90%) of LDL oxidation was observed. Inhibition was heat-resistant since heat-treated (60°C for 15 min) SAG still retains the activity, suggesting that enzymatic activity is not involved. Inhibitory activity was, however, completely or partially abolished by pretreatment of SAG with alkylating reagents NEM and p-hydroxy mercury benzoate (PHMB), respectively. The results indicated that free SH groups in SAG are the major contributors to this activity. Furthermore, metallothionein, a small metal binding protein consisting of 20 cysteine residues out of 61 amino acids (Nordberg & Kojima, 1979) showed a similar inhibitory curve as SAG. Glutathion (GSH), an additional cysteine containing peptide showed a 25% inhibition at a concentration of 100 µM. Inhibition of copper-induced LDL oxidation was, however, not observed in other known antioxidant enzymes such as superoxide dismutase, catalase or other proteins such as BSA, and cytochrome C. These results clearly showed that by binding and chelating copper ion through its free SH groups, SAG prevents copperinitiated free radical reactions leading to LDL oxidation and superoxide or hydrogen peroxide appear not to be involved in the process. To test whether SAG protection against LDL oxidation was solely mediated through copper binding, we initiated LDL oxidation by AAPH, a free radical generator. In this metal-ion free system, SAG also protects LDL oxidation (up to 85%) at a concentration of 59 µM (750 µg/mL). Thus, by metal binding and free radical scavenging, SAG acts as a protector against lipid peroxidation.

Example 26. SAG protects cytochrome C release and caspase activation induced by metal ions

Since cytochrome C release from mitochondria and caspase activation are the key events in apoptosis (Liu et al., 1996; Yang et al., 1997; Li et al., 1997; Hengartner, 1998, for review, see Mignotte & Vayssiere, 1998), the levels of cytochrome C released into cytoplasm and potential activation of caspase upon metal treatments were measured. Treatment of cells with ZnSO4 induces a time-dependent release of cytochrome C in cytoplasm. Compared to the vector control cell (D1-6), the SAG overexpressing cell (D12-1) has much less cytoplasmic release of cytochrome C. Likewise, activation of caspase 7, shown as disappearance of pro-enzyme form, was seen in a time-dependent manner post zinc treatment. More activation was seen in vector control cell (D1-6) than that in the SAG overexpressing

cell (D12-1). A similar result was obtained with CPP32 (caspase 3) activation. A significant difference, however, was not seen in cytochrome C release or caspase activation between D1-6 and D12-1 cells upon copper treatment. This is consistent with the lack of difference in morphological changes between the two lines upon copper treatment, although DNA fragmentation was obvious only in the vector control cells. To further examine potential protection of SAG against metal-induced cytochrome C release and CPP32 activation, cytochrome C release and CPP32 activation was measured in 293 cells transiently transfected with SAG expressing plasmid followed by exposure to copper. A significant amount of cytochrome C started to release 6 hours post CuSO₄ (2.0 mM) treatment and lasted up to 12 hours. Expression of SAG delayed cytochrome C release for up to 16 hours. Activation of caspase 7 was seen in the vector control cells 12 hours and 16 hours post copper treatment. No significant activation was seen in SAG transfectants. The similar result was seen with CPP32 antibody. For zinc treatment, no difference was detected in cytochrome C release and caspase activation between control cells and SAG transfectants, consistent with the lack of difference in morphological signs of apoptosis. These results indicate that metal treatment induces cytochrome C release and caspase activation during apoptosis which can be largely prevented or delayed by SAG and there is a good correlation between morphological signs of apoptosis and cytochrome C release/caspase activation.

Example 27: SAG protects against neuronal apoptosis

5

10

15

20

25

30

SAG was transfected into HY5Y human neuroblastoma cells and a few stable lines were selected which expressed exogenous SAG as determined by Western blot. One SAG-transfectant (SYW-20) and a vector control (SYV-3) were used to determine their sensitivity to metal ions, zinc and copper. Treatment with 1.25 mM CuSO₄ or 200 µM ZnSO₄ for 16 hours induced cell shrinkage and detachment in the neo control cells, but to a less extent in SAG-expressing cells. The morphological difference was more obviously seen with the zinc treatment. To determine the nature of cell death, we performed TUNEL assay, a fluorescein labelling assay of free 3'-OH termini generated from cleavage of genomic DNA during apoptosis.

In Situ cell death assay (TUNEL assay) was performed according to the manufacturer's instructions (Boehringer Mannheim). Briefly, 5 x 10⁴ cells were plated into the 8-well glass slides. After treatment with 1.25 mM copper (CuSO₄) or 200 μM zinc (ZnSO₄) for 16 hours, cells were fixed with 0.5% glutaraldehyde for 10 min, then washed

with PBS twice. The fixed cells were incubated in permeabilization solution (0.1% Triton X-100, 0.1% sodium citrate) for 2 min on ice. The TUNEL reaction mixture (50 μL) was added to samples and incubated for 1 hour at 37°C followed by 3 times wash with PBS. Samples were embedded with antifade prior to analysis under a fluorescence microscope. Substantially more fluorescein staining was seen in the vector control cells after 16 hours treatment with 1.25 mM CuSO₄, or 200 μM ZnSO₄. The results indicate that expression of SAG protects neuronal cells from apoptosis.

Example 28. SAG stimulates proliferation

5

10

15

20

25

30

To test potential growth stimulation activity, SAG RNA (8 μ g/mL or 25 μ g/mL), along with the control β -galactosidase (25 μ g/mL), was injected into serum-starved NIH 3T3 fibroblast monolayer. Approximately 50 cells attached to the glass coverslip within an etched circle were injected. A 3-hour pulse of [³H]thymidine (5 μ Ci/mL, Amersham) was performed 10 to 24 hours after injection. Cultures were washed with isotonic phosphate-buffered saline and fixed in 3.7% (vol/vol) formaldehyde. Induction of [³H]thymidine incorporation (an indicator of DNA synthesis) into the nuclei of serum-starved fibroblast cells was obviously observed in SAG-injected cells. In contrast, injection of β -galactosidase does not induce DNA synthesis and no [³H]thymidine incorporation was observed. The results clearly indicate that human SAG has proliferative activity to stimulate cell growth.

Growth promotion activity of SAG was also examined in human neuroblastoma cells (SY5Y), overexpressing hSAG protein by hSAG cDNA transfection. Both the vector-expressing control cells and SAG overexpressing cells were first serum-starved for 48 hours, followed by 3H-thymidine labelling for 16 hours in either serum-starved or 1% serum conditions. Cells were washed, lysed and counted in a liquid scintillation counter for 3H, an assay for the measurement of 3H-thymidine incorporation into DNA (S-phase entry). Compared to the vector control cells, SAG-expressing cells have 10-fold more 3H-thymidine incorporation in both conditions (serum-free or 1% serum), indicating that SAG stimulates cell proliferation/growth.

Growth promotion activity of SAG was also examined in yeast. As described in Example 22, the yeast homolog of human SAG gene is essential for yeast growth. To correlate yeast growth rate with SAG expression, hSAG expressing plasmid was constructed under control of Gal promoter. The plasmid was transformed into heterozygous ySAG knockout and transformants were sporulated and dissected. Haploid ySAG knockout clone

that contained hSAG plasmid was identified and analyzed. In the uninduced condition, little SAG expression due to the leakness of the promoter led to formation a tiny clone compared to the full size wildtype clone. Under induced condition, SAG expression level increased and clone size also increased. This experiment clearly demonstrated that SAG promotes cell growth in a dose-dependent manner.

5

10

It is to be understood that the invention is not to be limited to the exact details of operation, or to the exact compounds, compositions, methods, procedures or embodiments shown and described, as obvious modifications and equivalents will be apparent to one skilled in the art, and the invention is therefore to be limited only by the full scope of the appended claims.

I claim

20

1. An isolated and purified DNA sequence substantially similar to the DNA sequence shown in SEQ ID 1.

- An isolated and purified DNA sequence that hybridizes to the DNA sequence shown
- 5 in SEQ ID 1 under high stringency hybridization conditions.
 - 3. An isolated and purified DNA sequence that consists essentially of the DNA sequence shown in SEQ ID 1.
 - 4. A recombinant DNA molecule comprising the isolated and purified DNA sequence of Claim 1, 2, or 3 subcloned into an extra-chromosomal vector.
- A recombinant host cell comprising a host cell transfected with the recombinant DNA molecule of Claim 4.
 - 6. A recombinant host cell deposited with the ATCC under accession number 98402.
 - 7. An isolated and purified DNA sequence substantially similar to the DNA sequence shown in SEQ ID 3.
- 8. An isolated and purified DNA sequence that hybridizes to the DNA sequence shown in SEQ ID 3 under high stringency hybridization conditions.
 - An isolated and purified DNA sequence that consists essentially of the DNA sequence shown in SEQ ID 3.
 - 10. A recombinant DNA molecule comprising the isolated and purified DNA sequence of Claim 7, 8, or 9 subcloned into an extra-chromosomal vector.
 - 11. A recombinant host cell comprising a host cell transfected with the recombinant DNA molecule of Claim 10.
 - 12. A recombinant host cell deposited with the ATCC under accession number 98403.
 - 13. A recombinant host cell deposited with the ATCC under accession number 98404.
- 25 14. A recombinant host cell deposited with the ATCC under accession number 98405.
 - 15. An isolated and purified DNA sequence selected from the group consisting of SEQ ID 11, SEQ ID 13, SEQ ID 21, SEQ ID 23, SEQ ID 25, SEQ ID 27, SEQ ID 29, SEQ ID 31, SEQ ID 33, SEQ ID 35, SEQ ID 37, SEQ ID 39, SEQ ID 41, SEQ ID 43, SEQ ID 45, SEQ ID 47, and SEQ ID 49.
- 30 16. A recombinant DNA molecule comprising an isolated and purified DNA sequence of Claim 15, subcloned into an extra-chromosomal vector.
 - 17. A recombinant host cell comprising a host cell transfected with a recombinant DNA molecule of Claim 16.
 - 18. A substantially purified recombinant polypeptide, wherein the amino acid sequence of

the substantially purified recombinant polypeptide is substantially similar to the amino acid sequence shown in SEQ ID 2.

19. A substantially purified recombinant polypeptide, wherein the amino acid sequence of the substantially purified recombinant polypeptide consists essentially of the amino acid sequence shown in SEQ ID 2.

5

15

20

- 20. A substantially purified recombinant polypeptide, wherein the amino acid sequence of the substantially purified recombinant polypeptide is substantially similar to the amino acid sequence shown in SEO ID 4.
- 21. A substantially purified recombinant polypeptide, wherein the amino acid sequence of the substantially purified recombinant polypeptide consists essentially of the amino acid sequence shown in SEQ ID 4.
 - 22. A substantially purified recombinant polypeptide, wherein the amino acid sequence of the polypeptide is selected from the group consisting of SEQ ID 12, SEQ ID 14, SEQ ID 22, SEQ ID 24, SEQ ID 26, SEQ ID 28, SEQ ID 30, SEQ ID 32, SEQ ID 34, SEQ ID 36, SEQ ID 38, SEQ ID 40, SEQ ID 42, SEQ ID 44, SEQ ID 46, SEQ ID 48, and SEQ ID 50.
 - 23. An antibody that selectively binds polypeptides with an amino acid sequence substantially similar to the amino acid sequence of Claim 18, 19, 20, 21 or 22.
 - 24. A method of detecting SAG protein in cells, comprising contacting cells with the antibody of Claim 23 and incubating the cells in a manner that allows for detection of the SAG protein-antibody complex.
 - 25. A diagnostic assay for detecting cells containing SAG mutations, comprising isolating total genomic DNA from the cell and subjecting the genomic DNA to PCR amplification using primers derived from the isolated and purified DNA sequence of Claim 1, 2, 3, 7, 8, 9, or 15, and determining whether the resulting PCR product contains a mutation.
- 26. A diagnostic assay for detecting cells containing SAG mutations, comprising isolating total cell RNA, subjecting the RNA to reverse transcription-PCR amplification using primers derived from the isolated and purified DNA sequence of Claim 1, 2, 3, 7, 8, 9, or 15 and determining whether the resulting PCR product contains a mutation.
 - 27. A method of isolating RNA containing stretches of polyA or polyC residues, comprising
 - (a) contacting an RNA sample with SAG protein in RNA binding buffer in the presence of a reducing agent;
 - (b) incubating the RNA-SAG protein mixture with the antibody of Claim 23;
 - (c) isolating the antibody-SAG protein-RNA complexes; and

- (d) purifying the RNA away from the antibody-SAG protein complex.
- 28. A method of isolating RNA containing stretches of polyU residues, comprising
- (a) contacting an RNA sample with SAG protein in RNA binding buffer in the absence of reducing agents;
 - (b) incubating the RNA-SAG protein mixture with the antibody of Claim 23;
 - (c) isolating the antibody-SAG protein-RNA complexes; and
 - (d) purifying the RNA away from the antibody-SAG protein complex.
- 29. A method for isolating genes induced during cell apoptosis, comprising:
 - (a) treating one set of cells with OP and not treating a control set of cells;
- 10 (b) isolating RNA from each set of cells;

5

15

20

- (c) subjecting the RNA from each set of cells to the differential display procedure, wherein the RNA is reverse transcribed into cDNA and the cDNA is subjected to the polymerase chain reaction;
- (d) identifying cDNAs that are expressed in the OP-treated set of cells and not in the control set of cells; and
 - (e) cloning the OP-induced cDNAs.
 - 30. A method for protecting cells from apoptosis induced by redox reagents, comprising introducing into the cells an expression vector comprising the isolated and purified DNA sequence of Claim 1, 2, 3, 7, 8, 9, or 15, which is operatively linked to a DNA sequence that promotes the high level expression of the isolated and purified DNA sequence in the cells.
 - 31. A method for inhibiting the growth of tumor cells, comprising introducing into the tumor cells an expression vector comprising the isolated and purified DNA sequence of Claim 1, 2, 3, 7, 8, 9, or 15, which is operatively linked to a DNA sequence that promotes the high level expression of the antisense strand of the isolated and purified DNA sequence in the cells.
 - 32. A method for purifying SAG protein from bacterial cells comprising:
 - transfecting a bacterial host cell with a vector comprising the isolated and purified DNA sequence of Claim 1, 2, 3, 7, 8, 9, or 15 operatively linked to a promoter capable of directing gene expression in a bacterial host cell;
- b) inducing expression of the isolated and purified DNA sequence in the bacterial
 cells;
 - c) lysing the bacterial cells;
 - d) isolating bacterial inclusion bodies;
 - e) purifying SAG protein from the isolated inclusion bodies.

33. A pharmaceutical composition comprising the substantially purified recombinant polypeptide of Claim 18, 19, 20, 21, or 22 and a pharmaceutically acceptable carrier.

- 34. The pharmaceutical composition of Claim 33 wherein the substantially purified recombinant polypeptide comprises an oligomer.
- 5 35. A method of oxygen radical scavenging in an organism comprising administering an oxygen radical -reducing amount of the pharmaceutical composition of Claim 33 or 34 to the organism.
 - 36. A method of promoting the healing of a wound comprising administering the DNA sequence of Claim 1 to cells associated with the wound.
- 10 37. A method of promoting or inhibiting the growth of plant cells comprising administering the DNA sequence of Claim 1 or a DNA sequence which is complementary to the DNA sequence of Claim 1 to plant cells.

e e		

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT:
 - (A) NAME: Yi Sun
 - (B) STREET: 4841 Hillway Court
 - (C) CITY: Ann Arbor
 - (D) STATE: Michigan
 - (E) COUNTRY: USA
 - (F) POSTAL CODE (ZIP): 48105
 - (G) TELEPHONE: (313) 996-1959
 - (H) TELEFAX: (313) 996~7158
- (ii) TITLE OF INVENTION: Sensitive to Apoptosis Gene (SAG)
- (iii) NUMBER OF SEQUENCES: 50
- (iv) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)
- (2) INFORMATION FOR SEQ ID NO: 1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1140 base pairs
 - (B) TYPE: nucleic acid(C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 17..355
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION: 17..355
 - (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION:1..1140
 - (D) OTHER INFORMATION:/note= "Mouse SAG"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:
- GTTCTGCGCC GCCGCC ATG GCC GAC GTG GAG GAC GGC GAG GAA CCC TGC

 Met Ala Asp Val Glu Asp Gly Glu Glu Pro Cys

 1 5 10
- GTC CTT TCT TCG CAC TCC GGG AGC GCA GGC TCC AAG TCG GGA GGC GAC

 Val Leu Ser Ser His Ser Gly Ser Ala Gly Ser Lys Ser Gly Gly Asp

 15

 20

 25



W	O 99	/3251	4												PCT/US	98/26705
			TCT Ser													145
		GAG	TGC Cys													193
			CGA Arg													241
			GAG Glu													289
									Leu						GTA Val	337
			ATC Ile				GAGG	TGG	CCCA	GGCG	ст с	CTGG	TGTG	G		385
TTGO	CTGA	ccc	TGGA	CAAA	GA C	TAAA	CACT	G CA	GGGG	ATTC	ATC	CTTG	AGA	GAGA	GAGGAT	445
GCTC	STGC	GCC	TTTG	AGAC	TC A	CCAA	AGGC	T TG	CTTT	ATTA	ATT	TGTC	TGT	TTAG	TTTTGG	505
GAA	ATTC	тст	ACAA	TTAA	GA T	AATT	TGTT	'A AA	AATG	GCCT	TTC	CTAC	CTC	TGGT	GTGTGT	565
GTG?	rgat.	ACG	AATG	CATA	GA A	GAGC	GAGA	A CA	CCAG	AAAA	TGA	TCTI	TGT	TTAT	CTGTAC	625
CCA	CGAC	TGG	AACA	TTGT	GT I	CACA	GAAG	A AC	OTTA:	TTTG	TGT	PATT	GCT	TGAG	GGTTAA	685
AAA	ATAG	ATA	AACG	AATG	TT A	CAGI	AACA	ra aj	AAAA	TGCA	TTC	AAAA	GCC	GACI	CCTCCT	745
AAT	CCTT	TTT	GTGT	TGGG	AG A	GAGG	CAAG	C GA	AGGCC	CACCO	TGC	TGTC	TTC	ATTI	GCTGTG	805
AAT	GAGG	ATT	TTAA	CCTC	CA C	CTCAG	TGA	AG AG	GCG1	PAACT	GTC	GGG1	AAA	CTGT	PTATAA	865
GCG	TAAC	TGT	CGGG	TAA	CG C	CTTI	GTCI	rc ci	rgaci	TCTC	CAT	CTTT	GAC	TTGO	CCAGGA	925
AGC	CTGG	ATT	GTTC	CAACO	CAC 1	TAGT	rtct?	AA AG	BAACT	rgtti	TC	rgtti	rttg	CCG	AAGGTTG	985
TAT	TGTA	TGT	TTT	\GTC!	AAA 2	\ATA1	rtag:	ra Go	SAAA	ATGGC	TTI	ACTAC	STAT	AAC	ACTGAAG	1045
TTC	ATTA	ATGC	AATO	STTT	CAA !	LAAA1	ATAT!	rg to	GCTT'	rgag1	TA?	rtaa <i>i</i>	AGTT	TGA:	ГАТАТАС	1105

(2) INFORMATION FOR SEQ ID NO: 2:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear

TCTTAAAATC ATTAAACTAA TTCATCAATT AAATG

- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:



Met Ala Asp Val Glu Asp Gly Glu Glu Pro Cys Val Leu Ser Ser His 1 5 10 15

Ser Gly Ser Ala Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp
35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys
50 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys
65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 3:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..339
- (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION:1..339
- (ix) FEATURE:
 - (A) NAME/KEY: misc_feature
 - (B) LOCATION:1..754
 - (D) OTHER INFORMATION:/note= "Human SAG"
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

1 5 10 15

TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu

20 25 30

AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT 144



V	VO 99	/3251	4												PCT/	US98/26705
Lys	Lys	Trp 35	Asn	Ala	Val	Ala	Met 40	Trp	Ser	Trp	Asp	Val 45	Glu	Cys	Asp	
															TGT Cys	192
										GTG Val 75					TGT Cys 80	240
										CTG Leu					AAC Asn	288
														Ile	GGC Gly	336
AAA Lys		GAGT	GGT '	TAGA	AGGC'	TT C	TTAG	CGCA	G TT	GTTC:	AGAG	CCC	TGGT	GGA		389
TCT	TGTA	ATC	CAGT	GCCC	TA C	AAAG	GCTA	G AA	CACT	ACAG	GGG	ATGA	ATT	CTTC	AAATAG	449
GAG	CCGA	TGG	ATCT	GTGG	тс т	TTGG	ACTC	A TC	AAAG	CCTT	GGT	TAGC	ATT	TGTC	AGTTT	r 509
ATC	TTCA	GAA	ATTC	TCTG	TG A	TTAA	GAAG	а та	ATTT	ATTA	AAG	GTGG	TCC	TTCC	TACCT	569
TGT	GGTG	TGT	GTCG	CGCA	CA C	AGCT	TAGA	A GT	GCTA	TAAA	AAA	GGAA	AGA	GCTC	CAAAT'	r 629
GAA	TCAC	CTT	ATAA	TTTA	.cc c	ATTT	CTAT	A CA	ACAG	GCAG	TGG	AAGC	AGT	TTCG	AGACT'	r 689
ттт	CGAT	GCT	TATG	GTTG	AT C	AGTT	AAAA	A AG	AATG	TTAC	AGT	AACA	AAT	AAAG	TGCAG	r 749
TTA	AA															754

(2) INFORMATION FOR SEQ ID NO: 4:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60



Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys
65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110

Lys

- (2) INFORMATION FOR SEO ID NO: 5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5:

AAGCTTTTTT TTTTTTR

18

- (2) INFORMATION FOR SEQ ID NO: 6:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 13 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 (A) DESCRIPTION: /desc = "Oligonucleotide: P2
 upstream primer"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

AAGCTTNNNN NNN

- (2) INFORMATION FOR SEQ ID NO: 7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 25 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "Oligonucleotide SAG TA.01"



(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:	
CGGGATCCCC ATGCCCGACG TGAGG	25
(2) INFORMATION FOR SEQ ID NO: 8:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "oligonucleotide SAG T.02"</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:	
CGGGATCCTC ATTTGCCGAT TCTTTG	26
(2) INFORMATION FOR SEQ ID NO: 9:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "oligonucleotide P.01"</pre>	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:	
TATGGCTAGC ATGGCCGACG TGGAGG	26
(2) INFORMATION FOR SEQ ID NO: 10:	
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: peptide	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 10:	
Gln Asn Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg	



121	INFORMATION	FOR	SEO	TD	NO:	11:

	CHARTON	CHARACTERISTICS:
(1)	SECUENCE	CHARACTERISTICS:

- (A) LENGTH: 747 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

- (A) NAME/KEY: CDS
- (B) LOCATION: 1..270

(ix) FEATURE:

- (A) NAME/KEY: mat_peptide
- (B) LOCATION: 1..270

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

Met									ACC Thr							48
									GGC					TCC		96
Ser	Gly	Ser	Ser 20	GIĀ	ser	ьуs	ser	25	GIA	ASP	ьys	Mec	30	ser	Leu	
															GAT Asp	144
												Asp			CTG Leu	192
	Thr					Val					Glu				ATT Ile 80	240
					Ala				GGG Gly		AACA	.GAA	CAAT	CGCT	GC	290
CCI	стст	GCC	AGCA	GGAC	TG G	GTGG	TCCA	A AG	AATC	GGCA	raa .	GAGA	GTG	GTTA	GAAGGC	350
TTC	TTAG	CGC	AGTT	GTTC	AG A	GCCC	TGGT	G GA	ATCTI	GTAA	TCC	AGTO	CCC	TACA	AAGGCT	410
AGA	ACAC	TAC	AGG	GATO	AA 1	TTCTT	CAA	AT AC	GAGC	CGAT	GGF	ATCTO	STGG	TCTT	TGGACT	470
CAT	CAA	AGCC	TTG	TTA(GCA T	rttgi	CAG	ייד ידיו	ratci	TCAC	AAA	ATTC	CTG	TGAT	TAAGAA	530
GA?	'AAT'	ТАТ	TAA	AGGT	GT (CTTC	CTA	C TO	CTGT	GTGT	r GTY	STCG	CGCA	CAC	AGCTTAG	590
AAC	TGC:	rata	AAA	AAGG	AAA (GAGC'	CCA	AA T	rgaa:	CAC	TTI	ATAA'	PTTA	CCC	ATTTCTA	650
тас	TAAC	AGGC	AGT	GGAA	GCA (GTTT	CGAG	AC T	TTTT(GAT	G CT	ratg	GTTG	ATC	AGTTAAA	710



- (2) INFORMATION FOR SEQ ID NO: 12:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 90 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Met Pro Val Leu Asp Val Lys Leu
50 55 60

Lys Thr Asn Lys Arg Thr Val Leu Trp Ser Gly Glu Asn Val Ile Ile 65 70 75 80

Pro Ser Thr Thr Ala Ala Cys Pro Cys Gly 85 90

- (2) INFORMATION FOR SEQ ID NO: 13:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 706 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION:1..291
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION: 1...291
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

1 5 10 15

TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

	go.	
•		•

			AAC Asn												GAT Asp	144
															TGT Cys	192
						Cys									AAC Asn 80	240
					Cys										GJY GGC	288
AAA Lys	TGA	GAGT	GGT	TAGA	AGGC	тт с	TTAG	CGCA	G TT	GTTC.	AGAG	ccc	TGGT	GGA		341
TCT	TGTA	ATC	CAGT	GCCC	TA C	AAAG	GCTA	G AA	CACT	ACAG	GGG	ATGA	TTA	CTTC	AAATAG	401
GAG	CCGA	TGG	ATCT	GTGG	TC I	TTGG	ACTO	A TO	AAAG	CCTT	GGT	TAGO	ATT	TGTC	AGTTTT	461
ATC	TTCA	GAA	ATTC	TCTG	TG A	AATT.	GAAG	A TA	ATTI	ATTA	AAG	GTGG	TCC	TTCC	TACCTC	521
TGT	GGTG	TGT	GTCG	CGC	CA C	CAGCI	TAGA	A GI	GCTA	TAAA	AAA	.GGA.	AGA	GCTC	CAAATT	581
GAA	TCAC	CTT	ATAA	\TTT#	ACC (CATT	CTAT	TA CA	ACAC	GCAG	TGG	AAGO	CAGT	TTC	AGACTT	641
ттт	CGAT	GCT	TATO	GTTC	GAT (CAGT	TAAA?	AA AC	TAAE	STTAC	AG1	TAAC	LAAT	AAAC	STGCAGT	701
TTF	AA							•						,		706

(2) INFORMATION FOR SEQ ID NO: 14:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 97 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp $35 \hspace{1cm} 40 \hspace{1cm} 45$

Thr Cys Ala Ile Cys Arg Val Gln Val Met Val Val Trp Gly Glu Cys 50 60

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 65 70 75 80



Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 85 90 95

Lys

- (2) INFORMATION FOR SEQ ID NO: 15:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 18 base pairs
 - (B) TYPE: nucleic acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "oligonucleotide hSAG. M1"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

GCCATCTGCA GGGTCCAG

18

- (2) INFORMATION FOR SEQ ID NO: 16:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 27 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = "oligonucleotide SAG T.02L"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

GGATCCTCAT TTGCCGATTC TTTGGAC

27

- (2) INFORMATION FOR SEQ ID NO: 17:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 58 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 - (A) DESCRIPTION: /desc = *oligonucleotide
 SAGKanMX4-5*
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

TTCTCCAGTG GCAGAGACT TTAAAGAGAA ATAGTTCAAC CGTACGCTGC AGGTCGAC



(2)	INFORMATION FOR SEQ ID NO: 18:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 59 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
4	<pre>(ii) MOLECULE TYPE: other nucleic acid</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:	
ACC'	TCGGTAT GATTTAAATG TTTACGGGCA ATTCATTTTT ATCGATGAAT TCGAGCTCG	59
(2)	INFORMATION FOR SEQ ID NO: 19:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "oligonucleotide SAG pcr 5"	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:	
TTC	CTCCAGTG GCAGAGAAC	19
(2)	INFORMATION FOR SEQ ID NO: 20:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "oligonucleotide SAG pcr 3"</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:	
ATO	GATTTAAA TGTTTACGGG C	21
(2)) INFORMATION FOR SEQ ID NO: 21:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 754 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: double	



(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION:1..339

(ix) FEATURE:

(A) NAME/KEY: mat_peptide

(B) LOCATION:1..339

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

(AI) Bagomida Bassilla san ang ang ang	
ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15	48
TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC Ser Gly Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 25	96
AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35	144
ACG AGC GCC ATC TGC AGG GTC CAG GTG ATG GAT GCC TGT CTT AGA TGT Thr Ser Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60	192
CAA GCT GAA AAC AAA CAA GAG GAC TGT GTT GTG GTC TGG GGA GAA TGT Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80	240
AAT CAT TCC TTC CAC AAC TGC TGC ATG TCC CTG TGG GTG AAA CAG AAC Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95	288
AAT CGC TGC CCT CTC TGC CAG CAG GAC TGG GTG GTC CAA AGA ATC GGC Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110	336
AAA TGAGAGTGGT TAGAAGGCTT CTTAGCGCAG TTGTTCAGAG CCCTGGTGGA	389
TCTTGTAATC CAGTGCCCTA CAAAGGCTAG AACACTACAG GGGATGAATT CTTCAAATAG	449
GAGCCGATGG ATCTGTGGTC TTTGGACTCA TCAAAGCCTT GGTTAGCATT TGTCAGTTTT	509
ATCTTCAGAA ATTCTCTGTG ATTAAGAAGA TAATTTATTA AAGGTGGTCC TTCCTACCTC	569
TGTGGTGTGT GTCGCGCACA CAGCTTAGAA GTGCTATAAA AAAGGAAAGA GCTCCAAATT	629
GAATCACCTT ATAATTTACC CATTTCTATA CAACAGGCAG TGGAAGCAGT TTCGAGACTT	689
TTTCGATGCT TATGGTTGAT CAGTTAAAAA AGAATGTTAC AGTAACAAAT AAAGTGCAGT	749
TTAAA	754

		*	
ż			

- (2) INFORMATION FOR SEQ ID NO: 22:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp
35 40 45

Thr Ser Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys
65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110

Lys

- (2) INFORMATION FOR SEQ ID NO: 23:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..339
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION:1..339
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC
Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

1 5 10 15



		AGC Ser									_					96
		TGG Trp 35														144
		GCC Ala														192
		GAA Glu														240
		TCC Ser														288
				Leu										Ile	GGC Gly	336
AAA Lys		GAGT	GGT	TAGA	AGGC	тт с	TTAG	CGCA	g TT	GTTC	AGAG	ccc	TGGT	GGA		389
TCT	TGTA	ATC	CAGT	GCCC	TA C	AAAG	GCTA	G AA	CACT	ACAG	GGG	ATGA	ATT	CTTC	AAATAG	449
GAG	CCGA	TGG	ATCT	GTGG	TC T	TTGG	ACTC	A TC	AAAG	CCTT	GGT	TAGC	ATT	TGTC	AGTTTI	509
ATC	TTCA	GAA.	ATTC	TCTG	TG A	TTAA	GAAG	а та	ATTT	'ATTA	AAG	GTGG	TCC	TTCC	TACCTO	569
TGT	GGTG	TGT	GTCG	CGCA	CA C	AGCT	TAGA	A GT	GCTA	TAAA	. AAA	GGAA	AGA	GCTC	CAAATI	629
GAA	TCAC	CTT	ATAA	ATTT	.cc c	TTTA	CTAT	'A CA	ACAG	GCAG	TGG	AAGC	AGT	TTCG	AGACTT	689
TTI	CGAT	GCT	TATG	GTTG	AT C	AGTT	'AAAA	A AG	AATG	TTAC	AGT	AACA	TAA	AAAG	TGCAGI	749
TTA	AA															754

(2) INFORMATION FOR SEQ ID NO: 24:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30



Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Ser Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 25:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION:1..339
- (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide

70

- (B) LOCATION:1..339
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

												CTG Leu				48
												ATG Met				96
												GTG Val 45				144
		Ala										TGT Cys				192
CAA	GCT	GAA	AAC	AAA	CAA	GAG	GAC	TGT	GTT	GTG	GTC	TGG	GGA	GAA	TGT	240

AAT CAT TCC TTC CAC AAC TGC TGC ATG TCC CTG TGG GTG AAA CAG AAC 288

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Trp Gly Glu Cys



·· -			
Asn His Ser Phe His Asn C 85	Cys Cys Met Ser Leu 90	Trp Val Lys Gln Asn 95	
AAT CGC TGC CCT CTC TGC C Asn Arg Cys Pro Leu Cys C 100			336
AAA TGAGAGTGGT TAGAAGGCT. Lys	T CTTAGCGCAG TTGTTCA	GAG CCCTGGTGGA	389
TCTTGTAATC CAGTGCCCTA CA	AAGGCTAG AACACTACAG	GGGATGAATT CTTCAAATAG	449
GAGCCGATGG ATCTGTGGTC TT	TGGACTCA TCAAAGCCTT	GGTTAGCATT TGTCAGTTTT	509
ATCTTCAGAA ATTCTCTGTG AT	TAAGAAGA TAATTTATTA	AAGGTGGTCC TTCCTACCTC	569
TGTGGTGTGT GTCGCGCACA CA	GCTTAGAA GTGCTATAAA	AAAGGAAAGA GCTCCAAATT	629
GAATCACCTT ATAATTTACC CA	TTTCTATA CAACAGGCAG	TGGAAGCAGT TTCGAGACTT	689
TTTCGATGCT TATGGTTGAT CA	GTTAAAAA AGAATGTTAC	AGTAACAAAT AAAGTGCAGT	749
TTAAA			754

PCT/US98/26705

(2) INFORMATION FOR SEQ ID NO: 26:

WO 99/32514

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Ser Ala Ile Ser Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys
65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 27:



(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 754 base pairs
(B) TYPE: nucleic acid

				RAND: POLO				le									
	(ii)	MOL	ECUL	Е ТҮ	PE:	cDNA											
	(ix)	(A	•	: ME/K CATI			9										
	(ix)	(A		: ME/K CATI				ide									
	(xi)	SEÇ	UENC	E DE	SCRI	PTIC	N: S	EQ I	D NO	: 27	:						
				GAA Glu 5													48
				GGC Gly													96
				GCG Ala													144
				TGC Cys													192
				AAA Lys											TGT Cys 80		240
										Leu					AAC Asn		288
				Leu					Trp					Ile	GGC		336
AAA Lys		GAGT	GGT	TAGA	AGGC	TT C	TTAG	CGCA	G TT	GTTC	AGAG	ccc	TGGT	'GGA			389
тст	TGTA	ATC	CAGI	GCCC	TA C	AAAG	GCTA	G AA	CACI	ACAG	GGG	SATGA	TTA	CTTC	AAATA	G	449
GAG	CCG#	ATGG	ATCI	GTGG	TC I	TTGG	ACTO	A TO	CAAAC	CCTI	GG1	TAGO	ATT	TGTC	CAGTTT	T	509
ATC	TTC	AGAA	ATT	TCTC	TG A	ATTA?	GAAC	SA TA	\ATT1	TATTA	AA A	GTGC	TCC	TTCC	TACCT	С	569
TGT	rggto	STGT	GTC	GCGC	CA C	CAGCT	TAG	AA G	rgct <i>i</i>	AAATA	A AAJ	AGGA.	AAGA	GCT	CAAAT	т	629

		2.
÷		

GAATCACCTT ATAATTTACC CATTTCTATA CAACAGGCAG TGGAAGCAGT TTCGAGACTT

689

TTTCGATGCT TATGGTTGAT CAGTTAAAAA AGAATGTTAC AGTAACAAAT AAAGTGCAGT

749

TTAAA

754

- (2) INFORMATION FOR SEQ ID NO: 28:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Ser Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys
65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110

Lys

- (2) INFORMATION FOR SEO ID NO: 29:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION:1..339
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION:1..339

,			
	ę		

(xi)	SECUENCE	DESCRIPTION:	SEO	TΩ	NO ·	29.

													GCC Ala			48	3
													TTC Phe 30			9	6
													GAG Glu			14	4
													CTT Leu			19	2
						Glu					Val		GGA Gly			24	0
													AAA Lys		AAC Asn	28	8
									Trp				AGA Arg 110	Ile	GGC Gly	33	6
AAA Lys		GAGT	GGT	TAGA	AGGC	тт с	TTAG	CGCA	G TT	GTTC	AGAG	ccc	TGGT	GGA		38	19
TCT	TGTA	ATC	CAGT	GCCC	TA C	AAAG	GCTA	G AA	.CACT	ACAG	GGG	ATGA	ATT	CŤTC	AAATAG	G 44	19
GAG	CCGA	TGG	ATCT	GTGG	тс т	TTGG	ACTO	A TC	AAAG	CCTI	GGT	TAGO	ATT	TGTC	AGTTTT	50	9
ATC	TTCA	GAA	ATTC	TCTG	TG A	AATT	GAAG	A TA	PTTA	ATTA	AAG	GTGG	TCC	TTCC	TACCTO	5 6	59
TGT	GGTG	TGT	GTCG	CGCA	CA C	AGCT	TAGA	A GT	GCTA	TAAA	AAA	GGAA	AGA	GCTC	CAAATI	62	29
GAA	TCAC	CTT	ATAA	TTTA	cc c	TTTA	CTAT	'A CA	ACAG	GCAG	TGG	AAGC	AGT	TTCG	AGACTI	r 68	39
TTI	'CGA'I	GCT	TATO	GTTG	AT C	AGTI	'AAAA'	A AC	AATO	TTAC	AG1	AACA	AAT	AAAG	TGCAGT	r 74	49
TTA	AA															79	54

(2) INFORMATION FOR SEQ ID NO: 30:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 10

				;
	,			
		÷		
				ti
B				

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Trp Gly Glu Ser Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 105 100 Lys (2) INFORMATION FOR SEQ ID NO: 31: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 754 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear (ii) MOLECULE TYPE: cDNA (ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 1..339 (ix) FEATURE: (A) NAME/KEY: mat_peptide (B) LOCATION: 1..339 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31: ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC 48 Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 10 TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC 96 Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT 144 Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 ACG TGC GCC ATC TGC AGG GTC CAG GTG ATG GAT GCC TGT CTT AGA TGT 192 Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55

CAA GCT GAA AAC AAA CAA GAG GAC TGT GTT GTG GTC TGG GGA GAA TGT

60

		ń
çî.		
	, T	

W	O 99/	32514	i												PCT/U	S98/26705
Gln 2 65	Ala	Glu	Asn	Lys	Gln 70	Glu	Asp	Суз	Val	Val 75	Val	Trp	Gly	Glu	Суs 80	
AAT A	AAA Lys	TCC Ser	TTC Phe	CAC His 85	AAC Asn	TGC Cys	TGC Cys	ATG Met	TCC Ser 90	CTG Leu	TGG Trp	GTG Val	AAA Lys	CAG Gln 95	AAC Asn	288
								GAC Asp 105								336
AAA Lys	TGAG	SAGTO	GT 1	raga <i>i</i>	AGGC'	rr cr	PTAG(CGCAC	TTC	GTTC2	AGAG	ccc	rggtv	GGA		389
тстт	GTA.	ATC (CAGTO	ccc	ra c	AAAG	CTA	G AAC	CACT	ACAG	GGG	ATGA	ATT (CTTC	AAATAG	449
GAGC	CGAI	rgg z	ATCTO	TGG:	rc T	rtgg/	ACTC	A TC	AAAG	CCTT	GGT	rage:	ATT '	TGTC	AGTTTT	509
ATCT	TCAC	SAA A	ATTC	rctg:	rg a'	TAAG	GAAG	A TA	ATTT	ATTA	AAG	GTGG'	rcc '	TTCC	PACCTC	569
TGTG	GTGT	CT (STCGO	CGCA	CA C	AGCT	PAGA	A GTO	CTA	TAAA	AAA	GGAA.	AGA (GCTC	CAAATT	629
GAAT	CACC	TT A	ATAA?	rtta:	cc c	ATTT	CTAT	A CA	ACAG	GCAG	TGG	AAGC.	AGT '	TTCG.	AGACTT	689
TTTC	GATO	CT T	ratgo	STTG	AT C	AGTT	AAAA	A AG	AATG'	TTAC	AGT	AACA	AAT .	AAAG'	TGCAGT	749
TTAA	A															754
(2)		(i)		ENCE ENGT: YPE:	CHA H: 1 ami	RACT 13 ai no a	ERIS mino cid	32: TICS aci								
			LECU! QUEN					SEQ	ID N	0: 3	2:					
Met 1	Ala	Asp	Val	Glu 5		Gly	Glu	Glu	Thr 10		Ala	Leu	Ala	Ser 15	His	
Ser	Gly	Ser	Ser 20	Gly	Ser	Lys	Ser	Gly 25		Asp	Lys	Met	Phe 30		Leu	
Lys	Lys	Trp 35		Ala	Val	Ala	Met 40		Ser	Trp	Asp	Val 45		Cys	Asp	
Thr	Cys 50		Ile	Cys	Arg		Gln			Asp			Leu	Arg	Cys	

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys

Asn Lys Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly

. • Lys

		•												
(2)	INFO	RMAT:	ION 1	FOR S	SEQ I	D NO): 33	3:						
	(i)	(A (B (C) LEI) TY:) ST:	E CHANGTH: PE: 1 RANDI POLOG	: 754 nucle EDNE	l bas eic a SS: c	se pa acid doub	airs						
	(ii)	MOL	ECUL	E TY	PE:	cDNA								
	(ix)) NA	: ME/K CATI			9							
	(ix)		AN (: ME/K CATI				ide						
	(xi)	SEÇ	OUENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	: 33	:			
				GAA Glu 5										48
				GGC Gly										96
				GCG Ala										144
		Ala		TGC Cys										192
	Ala			AAA Lys										240
				AAG Lys 85									Asn	288
													GGC	336

105

TCTTGTAATC CAGTGCCCTA CAAAGGCTAG AACACTACAG GGGATGAATT CTTCAAATAG

GAGCCGATGG ATCTGTGGTC TTTGGACTCA TCAAAGCCTT GGTTAGCATT TGTCAGTTTT

AAA TGAGAGTGGT TAGAAGGCTT CTTAGCGCAG TTGTTCAGAG CCCTGGTGGA

100

Lys

22		
22		

389

•					
4					
4					
*					
					•
				3	
	£				
		•			
			A_{j}		
•					

WO 99/32514	1				PCT/U	S98/26705
ATCTTCAGAA A	ATTCTCTGTG	ATTAAGAAGA	TAATTTATTA	AAGGTGGTCC	TTCCTACCTC	569
TGTGGTGTGT C	GTCGCGCACA	CAGCTTAGAA	GTGCTATAAA	AAAGGAAAGA	GCTCCAAATT	629
GAATCACCTT A	ATAATTTACC	САТТТСТАТА	CAACAGGCAG	TGGAAGCAGT	TTCGAGACTT	689
TTTCGATGCT T	TATGGTTGAT	CAGTTAAAAA	AGAATGTTAC	AGTAACAAAT	AAAGTGCAGT	749
ТТААА						754
(3	SEQUENCE CI A) LENGTH:	HARACTERIST	ics:			
(1	B) TYPE: ar	mino acid				

(ii) MOLECULE TYPE: protein

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys
65 70 75 80

Asn His Ser Phe Lys Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 35:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS '
 - (B) LOCATION: 1...339
- (ix) FEATURE:



(A) NAME/KEY: mat_peptide

(B) LOCATION:1..339

(xi)	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:	35:
------	----------	--------------	-----	----	-----	-----

										TGC Cys						48
										GAC Asp						96
										TGG Trp						144
															TGT Cys	192
	Ala														TGT Cys 80	240
					Asn					Leu					AAC Asn	288
				Leu					Trp					Ile	GGC Gly	336
AAA Lys		GAGT	GGT	TAGA	AGGC	TT C	TTAC	CGCA	G TI	GTTC	AGAG	ccc	TGG1	GGA		389
TCT	TGTA	ATC	CAGI	GCCC	TA C	AAAC	GCTA	AG AA	CACI	ACAG	GGG	SATGA	LATT	CTTC	AAATAG	449
GAG	GCCG#	TGG	ATCI	GTG	STC 1	TTG	ACTO	CA TO	CAAAC	CCTT	GG1	TAGO	CATT	TGTC	AGTTTT	509
ATO	TTC	AGAA	ATTO	CTCTC	STG A	ATTA!	AGAAC	SA TA	ATTI	TATTA	AAC	GTGC	TCC	TTCC	TACCTC	569
TG	rggtc	TGT	GTC	GCGCZ	ACA (CAGC	rtag?	AA G	rgct <i>i</i>	XAAT	AA/	AGGAZ	AAGA	GCTC	CAAATT	629
GA	ATCAC	CTT	ATA	ATTT?	ACC (CATT	rcta:	ra c	AACAG	GCAC	TG	GAAG	CAGT	TTC	BAGACTT	689
TT?	rcga:	rgct	TATO	GTT	SAT (CAGT	raaa:	AA A	GAAT	STTAC	C AG	raac:	AAAT	AAAC	STGCAGT	749
TT	AAA															754

(2) INFORMATION FOR SEQ ID NO: 36:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein



(vi)	SECUENCE	DESCRIPTION:	SEO	TD	NO:	36:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp $35 \hspace{1cm} 40 \hspace{1cm} 45$

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Ser Met Ser Leu Trp Val Lys Gln Asn
85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 37:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1...339
- (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION:1..339

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 37:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

1 5 10 15

TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu

20
25
30

AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp

35 40 45

ACG TGC GCC ATC TGC AGG GTC CAG GTG ATG GAT GCC TGT CTT AGA TGT 192

			÷
	•,		

Thr	Cys 50	Ala	Ile	Суз	Arg	Val 55	Gln	Val	Met	Asp	Ala 60	Cys	Leu	Arg	Cys		
			AAC Asn												TGT Cys 80		240
															AAC Asn		288
				Leu										Ile	GGC		336
AAA Lys	TGA	GAGT	GGT '	TAGA	AGGC	тт с	TTAG	CGCA	G TT	GTTC.	AGAG	ccc	TGGT	GGA			389
TCT	TGTA	ATC	CAGT	GCCC	TA C	AAAG	GCTA	g aa	CACT	ACAG	GGG	ATGA	ATT	CTTC	AAATA	.G	449
GAG	CCGA	TGG	ATCT	GTGG	тс т	TTGG	ACTC	а тс	AAAG	CCTT	GGT	TAGC	ATT	TGTC	AGTTI	T.	509
ATC	TTCA	GAA	ATTC	TCTG	TG A	AATT.	GAAG	А ТА	ATTI	'ATTA	AAG	GTGG	TCC	TTCC	TACCI	rc.	569
TGT	GGTG	TGT	GTCG	CGCA	CA C	AGCI	TAGA	A GT	GCTA	AAAT	AAA	GGAA	AGA	GCTC	CAAAT	r	629
GAA	TCAC	CTT:	ATAA	TTTA	cc c	TTTA:	CTAT	'A CA	ACAC	GCAG	TGG	AAGC	AGT	TTCG	AGACT	T	689
TTI	CGAT	GCT	TATG	GTTG	AT C	AGTI	'AAAA'	A AG	AATC	TTAC	AGT	AACA	AAT	AAAG	TGCA	3T	749
TTA	.AA																754

(2) INFORMATION FOR SEQ ID NO: 38:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 38:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp $35 \hspace{1cm} 40 \hspace{1cm} 45$

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95



Asn Arg Ser Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110

Lys

(2) IN	FORMATION	FOR	SEO	ID	NO:	39:
--------	-----------	-----	-----	----	-----	-----

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION:1..339
- (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION: 1..339

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 39:

	 					GAG Glu									48
						TCG Ser									96
						ATG Met 40									144
						CAG Gln									192
						GAC Asp									240
				Asn		TGC Cys								Asn	288
			Leu					Trp					Ile	GGC Gly	336
AAA Lys	GAGT	GGT	TAGA	AGGC	тт с	TTAG	CGCA	G TT	GTTC	AGAG	ccc	TGGT	GGA		389

		rà.	
		,	
	¥		
		, .c	
			13.5

WO 99/325	PCT/US98	8/26705				
CTTGTAATC	CAGTGCCCTA	CAAAGGCTAG	AACACTACAG	GGGATGAATT	CTTCAAATAG	449

GAGCCGATGG ATCTGTGGTC TTTGGACTCA TCAAAGCCTT GGTTAGCATT TGTCAGTTTT 509

ATCTTCAGAA ATTCTCTGTG ATTAAGAAGA TAATTTATTA AAGGTGGTCC TTCCTACCTC 569

TGTGGTGTGT GTCGCGCACA CAGCTTAGAA GTGCTATAAA AAAGGAAAGA GCTCCAAATT 629

GAATCACCTT ATAATTTACC CATTTCTATA CAACAGGCAG TGGAAGCAGT TTCGAGACTT 689

TTTCGATGCT TATGGTTGAT CAGTTAAAAA AGAATGTTAC AGTAACAAAT AAAGTGCAGT 749

TTAAA 754

(2) INFORMATION FOR SEQ ID NO: 40:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys
50 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn Lys Ser Phe Lys Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 41:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: cDNA
- (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION:1..339

(ix) FEATURE:

- (A) NAME/KEY: mat_peptide
- (B) LOCATION:1..339

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15	
TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30	
AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45	
ACG TGC GCC ATC TGC AGG GTC CAG GTG ATG GAT GCC TGT CTT AGA AGT Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Ser 50 55 60	
CAA GCT GAA AAC AAA CAA GAG GAC TGT GTT GTG GTC TGG GGA GAA TGT Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80	1
AAT CAT TCC TTC CAC AAC TGC TGC ATG TCC CTG TGG GTG AAA CAG AAC Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95	
AAT CGC TGC CCT CTC TGC CAG CAG GAC TGG GTG GTC CAA AGA ATC GGC Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110	
AAA TGAGAGTGGT TAGAAGGCTT CTTAGCGCAG TTGTTCAGAG CCCTGGTGGA Lys	389
TCTTGTAATC CAGTGCCCTA CAAAGGCTAG AACACTACAG GGGATGAATT CTTCAAA	
GAGCCGATGG ATCTGTGGTC TTTGGACTCA TCAAAGCCTT GGTTAGCATT TGTCAGT	
ATCTTCAGAA ATTCTCTGTG ATTAAGAAGA TAATTTATTA AAGGTGGTCC TTCCTACC TGTGGTGTGT GTCGCGCACA CAGCTTAGAA GTGCTATAAA AAAGGAAAGA GCTCCAA	
GAATCACCTT ATAATTTACC CATTTCTATA CAACAGGCAG TGGAAGCAGT TTCGAGA	
TTTCGATGCT TATGGTTGAT CAGTTAAAAA AGAATGTTAC AGTAACAAAT AAAGTGC	
ттааа	754

(2) INFORMATION FOR SEQ ID NO: 42:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 113 amino acids



- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 42:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Ser 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

- (2) INFORMATION FOR SEQ ID NO: 43:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1...339
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION: 1...339
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 43:
- ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC

 Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

 1 5 10 15
- TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC

 Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu

 20 25 30
- AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT 144

		7	
	•		
÷			
			1.0
	4		
5			
14			

WO 99/32514 PCT/US	98/26705
Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45	
ACG TGC GCC ATC TGC AGG GTC CAG GTG ATG GAT GCC TGT CTT AGA TGT Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60	192
CAA GCT GAA AAC AAA CAA GAG GAC AGT GTT GTG GTC TGG GGA GAA TGT Gln Ala Glu Asn Lys Gln Glu Asp Ser Val Val Val Trp Gly Glu Cys 65 70 75 80	240
AAT CAT TCC TTC CAC AAC TGC TGC ATG TCC CTG TGG GTG AAA CAG AAC Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95	288
AAT CGC TGC CCT CTC TGC CAG CAG GAC TGG GTG GTC CAA AGA ATC GGC Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110	336
AAA TGAGAGTGGT TAGAAGGCTT CTTAGCGCAG TTGTTCAGAG CCCTGGTGGA Lys	389
TCTTGTAATC CAGTGCCCTA CAAAGGCTAG AACACTACAG GGGATGAATT CTTCAAATAG	449
GAGCCGATGG ATCTGTGGTC TTTGGACTCA TCAAAGCCTT GGTTAGCATT TGTCAGTTTT	509
ATCTTCAGAA ATTCTCTGTG ATTAAGAAGA TAATTTATTA AAGGTGGTCC TTCCTACCTC	569
TGTGGTGTGT GTCGCGCACA CAGCTTAGAA GTGCTATAAA AAAGGAAAGA GCTCCAAATT	629
GAATCACCTT ATAATTTACC CATTTCTATA CAACAGGCAG TGGAAGCAGT TTCGAGACTT	689
TTTCGATGCT TATGGTTGAT CAGTTAAAAA AGAATGTTAC AGTAACAAAT AAAGTGCAGT	749
TTAAA	754

(2) INFORMATION FOR SEQ ID NO: 44:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 44:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60



Gln Ala Glu Asn Lys Gln Glu Asp Ser Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 45:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: cDNA

- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..339
- (ix) FEATURE:

100

- (A) NAME/KEY: mat_peptide
- (B) LOCATION:1..339
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:

					GAC Asp											48
					TCC Ser											96
			-		GTG Val							-				144
		Ala			AGG Arg											192
	Ala				CAA Gln 70											240
					Asn					Leu					AAC Asn	288
AAT	CGC	TGC	CCT	CTC	TGC	CAG	CAG	GAC	TGG	GTG	GTC	CAA	AGA	ATC	GGC	336

110

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly

105



AAA TGAGAGT Lys	GGT TAGAAGG	CTT CTTAGCG	CAG TTGTTC#	AGAG CCCTGGT	ngga	389
TCTTGTAATC	CAGTGCCCTA	CAAAGGCTAG	AACACTACAG	GGGATGAATT	CTTCAAATAG	449
GAGCCGATGG	ATCTGTGGTC	TTTGGACTCA	TCAAAGCCTT	GGTTAGCATT	TGTCAGTTTT	509
ATCTTCAGAA	ATTCTCTGTG	ATTAAGAAGA	TAATTTATTA	AAGGTGGTCC	TTCCTACCTC	569
TGTGGTGTGT	GTCGCGCACA	CAGCTTAGAA	GTGCTATAAA	AAAGGAAAGA	GCTCCAAATT	629
GAATCACCTT	ATAATTTACC	CATTTCTATA	CAACAGGCAG	TGGAAGCAGT	TTCGAGACTT	689
TTTCGATGCT	TATGGTTGAT	CAGTTAAAAA	AGAATGTTAC	AGTAACAAAT	AAAGTGCAGT	749
TTAAA						754

(2) INFORMATION FOR SEQ ID NO: 46:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Ser Leu Arg Ser 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

(2) INFORMATION FOR SEQ ID NO: 47:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear

	•			
		13/1		
		p to		
	•		•	



(ii) MOLECULE TYPE: cDNA

(ix) FEATURE:

(A) NAME/KEY: CDS
(B) LOCATION:1..339

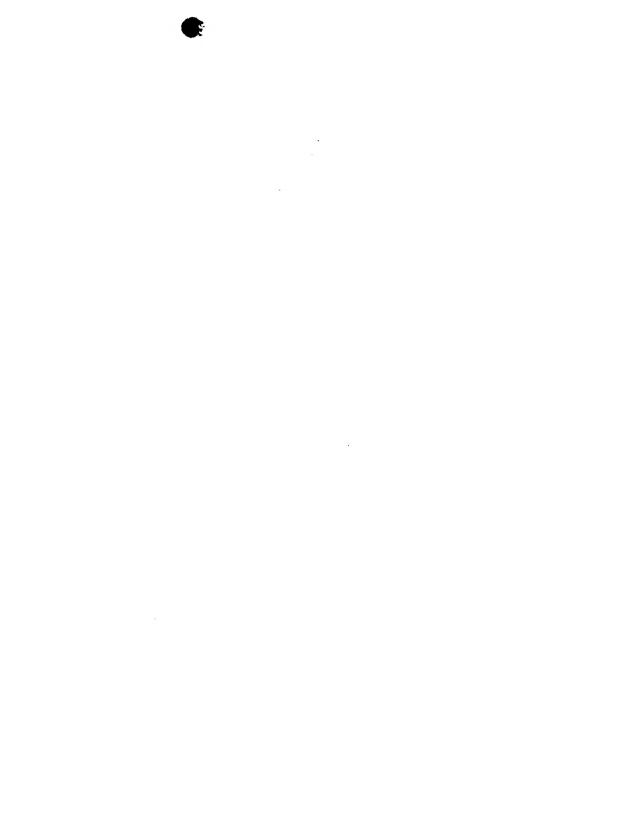
(ix) FEATURE:

(A) NAME/KEY: mat_peptide

(B) LOCATION:1..339

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 47:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15	48
TCC GGG AGC TCA GGC TCC AAG TCG GGA GGC GAC AAG ATG TTC TCC CTC Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30	96
AAG AAG TGG AAC GCG GTG GCC ATG TGG AGC TGG GAC GTG GAG TGC GAT Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45	144
ACG TGC GCC ATC TGC AGG GTC CAG GTG ATG GAT GCC TGT CTT AGA TGT Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60	192
CAA GCT GAA AAC AAA CAA GAG GAC TGT GTT GTG GTC TGG GGA GAA TGT Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Trp Gly Glu Cys 65 70 75 80	240
AAT CAT TCC TTC CAC AAC TGC TGC ATG TCC CTG TGG GTG AAA CAG AAC Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95	288
AAT CGC AGC CCT CTC AGC CAG CAG GAC TGG GTG GTC CAA AGA ATC GGC Asn Arg Ser Pro Leu Ser Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110	336
AAA TGAGAGTGGT TAGAAGGCTT CTTAGCGCAG TTGTTCAGAG CCCTGGTGGA Lys	389
TCTTGTAATC CAGTGCCCTA CAAAGGCTAG AACACTACAG GGGATGAATT CTTCAAATAG	449
GAGCCGATGG ATCTGTGGTC TTTGGACTCA TCAAAGCCTT GGTTAGCATT TGTCAGTTTT	509
ATCTTCAGAA ATTCTCTGTG ATTAAGAAGA TAATTTATTA AAGGTGGTCC TTCCTACCTC	569
TGTGGTGTGT GTCGCGCACA CAGCTTAGAA GTGCTATAAA AAAGGAAAGA GCTCCAAATT	629
GAATCACCTT ATAATTTACC CATTTCTATA CAACAGGCAG TGGAAGCAGT TTCGAGACTT	689
TTTCGATGCT TATGGTTGAT CAGTTAAAAA AGAATGTTAC AGTAACAAAT AAAGTGCAGT	749
TTAAA	754



- (2) INFORMATION FOR SEQ ID NO: 48:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 48:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Cys Asp 35 40 45

Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 55 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Ser Pro Leu Ser Gln Gln Asp Trp Val Val Gln Arg Ile Gly
100 105 110

Lys

- (2) INFORMATION FOR SEQ ID NO: 49:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 754 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: cDNA
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..339
 - (ix) FEATURE:
 - (A) NAME/KEY: mat_peptide
 - (B) LOCATION:1..339
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 49:

ATG GCC GAC GTG GAA GAC GGA GAG GAA ACC TGC GCC CTG GCC TCT CAC
Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His

1 5 10 15



W	O 99/	32514	1												PCT/	US98/26705
TCC Ser	GGG Gly	AGC Ser	TCA Ser 20	GGC Gly	TCC Ser	AAG Lys	TCG Ser	GGA Gly 25	GGC Gly	GAC Asp	AAG Lys	ATG Met	TTC Phe 30	TCC Ser	CTC Leu	96
														AGC Ser		144
														AGA Arg		192
														GAA Glu	TGT Cys 80	240
AAT Asn	CAT His	TCC Ser	TTC Phe	CAC His 85	AAC Asn	TGC Cys	TGC Cys	ATG Met	TCC Ser 90	CTG Leu	TGG Trp	GTG Val	AAA Lys	CAG Gln 95	AAC Asn	288
AAT Asn	CGC Arg	TGC Cys	CCT Pro 100	CTC Leu	TGC Cys	CAG Gln	CAG Gln	GAC Asp 105	TGG Trp	GTG Val	GTC Val	CAA Gln	AGA Arg 110	ATC Ile	GGC Gly	336
AAA Lys	TGAG	GAGTY	GGT '	TAGA	AGGC'	rt C	rtag(CGCA	G TT	GTTC:	AGAG	CCC	IGGT	GGA		389
TCT	rgta.	ATC (CAGT	GCCC'	ra c	AAAG	GCTA	G AA	CACT	ACAG	GGG.	ATGA.	ATT	CTTC	AAATA	3 449
GAG	CCGA'	rgg .	ATCT	GTGG'	rc T	TTGG.	ACTC.	A TC	AAAG	CCTT	GGT	TAGC	ATT	TGTC	AGTTT'	r 509
ATC'	PTCAG	GAA .	ATTC	TCTG	rg A	TTAA	GAAG.	A TA	ATTT.	ATTA	AAG	GTGG	TCC	TTCC'	TACCT	569
TGT	GGTG'	TGT	GTCG	CGCA	CA C	AGCT	TAGA	A GT	GCTA	TAAA	AAA	GGAA	AGA	GCTC	CAAAT	r 629
GAA'	TCAC	CTT .	ATAA	TTTA	CC C	ATTT	СТАТ	A CA	ACAG	GCAG	TGG	AAGC.	AGT	TTCG.	AGACT'	r 689
		GCT	TATG	GTTG.	AT C	AGTT	AAAA	A AG	AATG	TTAC	AGT	AACA	AAT	AAAG	TGCAG'	г 749
TTA	AA															754

(2) INFORMATION FOR SEQ ID NO: 50:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 113 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 50:

Met Ala Asp Val Glu Asp Gly Glu Glu Thr Cys Ala Leu Ala Ser His 1 5 10 15

Ser Gly Ser Ser Gly Ser Lys Ser Gly Gly Asp Lys Met Phe Ser Leu 20 25 30

Lys Lys Trp Asn Ala Val Ala Met Trp Ser Trp Asp Val Glu Ser Asp 35 40 45



Thr Cys Ala Ile Cys Arg Val Gln Val Met Asp Ala Cys Leu Arg Cys 50 60

Gln Ala Glu Asn Lys Gln Glu Asp Cys Val Val Val Trp Gly Glu Cys 65 70 75 80

Asn His Ser Phe His Asn Cys Cys Met Ser Leu Trp Val Lys Gln Asn 85 90 95

Asn Arg Cys Pro Leu Cys Gln Gln Asp Trp Val Val Gln Arg Ile Gly 100 105 110

Lys

	0.5				
		1.5			
	4 0				
iş:					
				٥	

INTERNATIONAL SEARCH REPORT

tnt .tional Application No PCT/US 98/26705

A. CLASSIFI IPC 6	CATION OF SUBJECT MATTER C07K14/47 C12N15/12 C12N15/10 //C12N15/82	A61K38/00	C07K16/18	C12Q1/68	3
According to	International Patent Classification (IPC) or to both n	ational classification a	ind IPC		
B. FIELDS S	EARCHED	41 41 41 41 41	nhole)		
IPC 6	umentation searched (classification system followe CO7K				
	on searched other than minimum documentation to				ed .
Electronic da	ata base consulted during the international search (name of data base an	id, where practical, search	n (erms used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT				
Category *	Citation of document, with Indication, where appr	opriate, of the relevan	nt passages		Relevant to claim No.
X	HILLIER L. ET AL.: "Soa tumor NbHPA Homo sapiens 5' " EMBL DATABASE,17 May 199 XP002106007 HEIDELBERG, DE Accession Number: W3871	s cDNA clone 96 (1996-05-	9 304814		1-5, 7-11, 18-21, 25,26
X	MARRA M. ET AL.: "Soar NbME13.5 14.5 Mus muscu 401232 5'" EMBL DATABASE,18 July 1 XP002106008 HEIDELBERG, DE Accession Number: W9809		1-5, 7-11, 18-21, 25,26		
X F	uther documents are listed in the continuation of bo	»x C.	Patent family men	nbers are usted in	annex
"A" docu con "E" earlie filin "L" docu whi cita "O" docu	categories of cited documents: ment defining the general state of the art which is residered to be of particular relevance ar document but published on or after the internation of date ment which may throw doubts on priority claim(s) of this cited to establish the publication date of anoth ation or other special reason (as specified) ument referring to an oral disclosure, use, exhibition are means ument published prior to the international filing date or than the priority date claimed	nal •; rer •	r later document publish or priority date and no cited to understand th invention X' document of particular cannot be considered involve an inventive so a victoria document of particular cannot be considered document is combine ments, such combina in the art. & document member of	relevance; the cla novel or cannot be tap when the doct relevance; the cla it o involve an inve d with one or mor tition being obvious the same patent fa	irred invention e considered to ment is taken alone inmed invention e considered to ment is taken alone inmed invention nitive step when the e other such docu- to a person skilled amily
Date of t	the actual completion of the international search		Date of mailing of the	international sear	ch report
	8 July 1999		20/07/199	99	
Name a	nd mailing address of the ISA European Patent Office, P.B. 5818 Patentli NL - 2280 HV Rijswijk Tet. (+31-70) 340-2040, Tx. 31 651 epo nl, Fav. (+31-71) 340-3016		Authorized officer	G	

INTERNATIONAL SEARCH REPORT

Int. | Nonal Application No PCT/US 98/26705

		PC1/US 98/26/US
C.(Continue	Ition) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Т	DUAN H. ET AL.: "SAG, a novel zinc RING finger protein that protects cells from apoptosis induced by redox agents" MOLECULAR AND CELLULAR BIOLOGY, vol. 19, no. 4, 1999, pages 3145-3155, XP002106009 US the whole document	1-26,29, 30,32-35
X	SUN Y.: "Induction of glutathione	29
	synthetase by 1,10-phenantroline" FEBS LETTERS, vol. 408, no. 1, 1997, pages 16-20, XP002106010 AMSTERDAM NL cited in the application abstract	
X	SUN Y. ET AL.: "Activation of p53 transcriptional activity by 1,10-phenanthroline, a metal chelator and redox sensitive compound" ONCOGENE, vol. 14, no. 4, 1997, pages 385-393, XP002106011 cited in the application abstract	29

INTERNATIONAL SEARCH REPORT

international application No.

PCT/US 98/26705

Box I	Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This Inte	ernational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely: see FURTHER INFORMATION sheet PCT/ISA/210
2.	Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful international Search can be carried out, specifically:
3.	Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box I	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This Ir	nternational Searching Authority found multiple inventions in this international application, as follows:
1. [As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2. [As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. [As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4. [No required additional search fees were timely paid by the applicant. Consequently, this international Search Report Is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Rer	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

FURTHER INFORMATION CONTINUED FROM PCT/ISA/ 210

Continuation of Box I.1

Although claims 25 and 26 are directed to a diagnostic method practised on the human/animal body, the search has been carried out and based on the alleged effects of the compound/composition.

Insofar as claims 30, 35-37 may be said to relate to methods in vivo, i.e methods of treatment of the human/animal body, then objection arises under Art 17.2.a.1 PCT, therefore a search has been carried out partially and based on the alleged effects of the compound/composition.